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# (12) United States Patent Bayon et al.

# (54) BIOSYNTHETIC IMPLANT FOR SOFT TISSUE REPAIR

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#### (58) Field of Classification Search

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#### (56) References Cited

# U.S. PATENT DOCUMENTS

1,187,158 A	6/1916	McGinley
3,118,294 A	1/1964	Van Laethem
3,124,136 A	3/1964	Usher
3,276,448 A	10/1966	Kronenthal et al.
3,320,649 A	5/1967	Naimer
3,364,200 A	1/1968	Ashton et al.
	(Cont	inued)

#### FOREIGN PATENT DOCUMENTS

CA	1317836	5/1993
DE	195 44 162	4/1997

(Continued)

# OTHER PUBLICATIONS

Ellouali M, et al., "Antitumor activity of low molecular weight fucans extracted from brown seaweed Ascophyllum Nodosum", *Anticancer Res.* Nov.-Dec. 1993; 12(6A):2011-9.

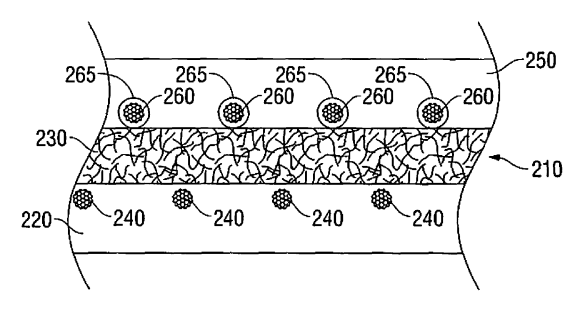
(Continued)

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#### (57) ABSTRACT

An implant and a process for preparing such an implant are disclosed. The implant includes a porous layer including collagen, a non-porous layer including a collagenic constituent, and a reinforcement component. The non-porous layer is joined to the porous layer and the reinforcement member is embedded into the non-porous layer. The porous layer has a three dimensional density ranging from about 20 mg collagen/cm<sup>3</sup> to about 200 mg collagen/cm<sup>3</sup>.

# 17 Claims, 5 Drawing Sheets



# US 9,242,026 B2 Page 2

(56)	Referen	ices Cited		5,543,441 A		Rhee et al.
TT	C DATENIT	DOCLIMENTS		5,565,210 A 5,567,806 A		Rosenthal et al. Abdul-Malak et al.
U.	S. PALENT	DOCUMENTS		5,569,273 A		Titone et al.
3,570,482 A	3/1071	Emoto et al.		RE35,399 E		Eisenberg
4,060,081 A		Yannas et al.		5,593,441 A		Lichtenstein et al.
4,173,131 A		Pendergrass et al.		5,595,621 A *		Light et al 156/80
4,193,137 A				5,601,571 A	2/1997	
4,248,064 A		Odham		5,607,590 A		Shimizu Bhas et al
4,294,241 A				5,614,587 A 5,618,551 A		Rhee et al. Tardy et al.
4,307,717 A		Hymes et al. Matsuda		5,639,796 A	6/1997	
4,338,800 A 4,476,697 A		Schäfer et al.		5,665,391 A	9/1997	
4,487,865 A		Balazs et al.		5,667,839 A	9/1997	
4,500,676 A		Balazs et al.		5,681,568 A		Goldin et al.
4,511,653 A	4/1985	Play et al.		5,686,115 A 5,690,675 A	11/1997	Vournakis et al. Sawyer et al.
4,527,404 A		Nakagaki et al.		5,697,978 A	12/1997	
4,591,501 A 4,597,762 A		Walter et al.		5,700,476 A		Rosenthal et al.
4,603,695 A		Ikada et al.		5,700,477 A		Rosenthal et al.
4,631,932 A		Sommers		5,709,934 A		Bell et al.
4,670,014 A		Huc et al.		5,716,409 A 5,720,981 A		Debbas Eisinger
4,709,562 A		Matsuda		5,732,572 A	3/1998	
4,748,078 A 4,759,354 A		Doi et al. Quarfoot		5,749,895 A		Sawyer et al.
4,769,038 A		Bendavid et al.		5,752,974 A		Rhee et al.
4,796,603 A		Dahlke et al.		5,766,631 A		Arnold
4,813,942 A		Alvarez		5,771,716 A		Schlussel
4,841,962 A		Berg et al.		5,785,983 A 5,800,541 A		Furlan et al. Rhee et al.
4,854,316 A 4,925,294 A		Davis Geshwind et al.		5,814,328 A		Gunasekaran
4,931,546 A		Tardy et al.		5,833,705 A		Ken et al.
4,942,875 A		Hlavacek et al.		5,861,034 A		Taira et al.
4,948,540 A	8/1990	Nigam		5,863,984 A		Doillon et al.
4,950,483 A		Ksander et al.		5,869,080 A 5,871,767 A		McGregor et al. Dionne et al.
4,970,298 A		Silver et al.		5,871,707 A 5,876,444 A	3/1999	
5,002,551 A 5,015,584 A		Linsky et al. Brysk		5,891,558 A		Bell et al.
5,147,374 A		Fernandez		5,899,909 A	5/1999	Claren et al.
5,162,430 A		Rhee et al.		5,906,937 A	5/1999	
5,171,273 A				5,911,731 A 5,919,232 A *	6/1999 7/1000	Pham et al. Chaffringeon et al 424/423
5,192,301 A 5,196,185 A		Kamiya et al. Silver et al.		5,919,232 A 5,919,233 A		Knopf et al.
5,190,183 A 5,201,764 A		Kelman et al.		5,922,026 A	7/1999	
5,206,028 A		Li	424/484	5,931,165 A		Reich et al.
5,217,493 A		Raad et al.		5,942,278 A		Hagedorn et al.
5,254,133 A				5,962,136 A 5,972,022 A	10/1999	Dewez et al.
5,256,418 A 5,263,983 A		Kemp et al. Yoshizato et al.		RE36,370 E	11/1999	
5,304,595 A		Rhee et al.		5,993,844 A	11/1999	Abraham et al.
5,306,500 A		Rhee et al.		5,994,325 A	11/1999	Roufa et al.
5,324,775 A	6/1994	Rhee et al.		5,997,895 A		Narotam et al.
5,328,955 A		Rhee et al.		6,001,895 A 6,008,292 A		Harvey et al. Lee et al.
5,334,527 A 5,339,657 A	8/1994 8/1004	Brysk McMurray		6,015,844 A		Harvey et al.
5,359,637 A 5,350,583 A		Yoshizato et al.		6,039,686 A		Kovac
5,356,432 A		Rutkow et al.		6,042,534 A		Gellman et al.
5,368,549 A		McVicker		6,042,592 A		Schmitt
5,376,375 A		Rhee et al.		6,043,089 A 6,051,425 A	3/2000	Sugiyama et al. Morota et al.
5,376,376 A 5,397,331 A		Himpens et al.		6,056,688 A		Benderev et al.
5,399,361 A		Song et al.		6,056,970 A		Greenawalt et al.
5,413,791 A		Rhee et al.		6,057,148 A		Sugiyama et al.
5,428,022 A	6/1995	Palefsky et al.		6,063,396 A	5/2000	Kelleher
5,433,996 A		Kranzler et al.		6,066,776 A 6,066,777 A		Goodwin et al. Benchetrit
5,441,491 A		Verschoor et al. Gazielly et al.		6,071,292 A		Makower et al.
5,441,508 A 5,456,693 A		Conston et al.		6,080,194 A		Pachence et al.
5,456,711 A				6,083,522 A		Chu et al.
5,466,462 A		Rosenthal et al.		6,120,539 A		Eldridge et al.
5,480,644 A				6,132,765 A		DiCosmo et al.
5,487,895 A 5,490,984 A		Dapper et al.		6,143,037 A 6,153,292 A		Goldstein et al. Bell et al.
5,512,291 A				6,165,488 A	12/2000	Tardy et al.
5,512,301 A		Song et al.		6,174,320 B1		Kugel et al.
5,514,181 A		Light et al.		6,179,872 B1	1/2001	Bell et al.
5,522,840 A		Krajicek		6,197,325 B1		MacPhee et al.
5,523,348 A		Rhee et al.		6,197,934 B1		DeVore et al.
5,536,656 A	//1996	Kemp et al.		6,197,935 B1	5/2001	Doillon et al.

# US 9,242,026 B2

Page 3

(56)	Referen	nces Cited	6,977,23 6,988,386			Matsuda Okawa et al.
U.S.	PATENT	DOCUMENTS	7,025,063	3 B2	4/2006	Snitkin et al.
			7,041,86	8 B2		Greene et al.
6,221,109 B1		Geistlich et al.	RE39,172 7,098,313			Bayon et al 424/444 Schaufler
6,224,616 B1 6,241,768 B1	5/2001 6/2001	Agarwal et al.	7,175,852			Simmoteit et al.
6,258,124 B1		Darois et al.	7,192,604			Brown et al.
6,262,332 B1		Ketharanathan	7,207,962 7,214,763			Anand et al. Ringeisen et al.
6,264,702 B1 6,277,397 B1		Ory et al. Shimizu	7,226,61			Yura et al.
6,287,316 B1		Agarwal et al.	7,229,453			Anderson et al.
6,306,424 B1		Vyakarnam et al.	7,931,69: 2001/0008930			Ringeisen 623/23.72 Tayot et al.
6,312,474 B1 6,328,686 B1	12/2001	Francis et al. Koyac	2002/009521			Carr, Jr. et al.
6,334,872 B1		Termin et al.	2002/0116070			Amara et al.
6,383,201 B1	5/2002		2003/0013989 2003/0023310			Obermiller et al. Brown et al.
6,391,333 B1 6,391,939 B2		Li et al. Tayot et al.	2003/0025310			Ringeisen
6,408,656 B1		Ory et al.	2003/0100954			Schuldt-Hempe
6,410,044 B1		Chudzik et al.	2003/011493′ 2003/013396′			Leatherbury et al. Ruszczak et al.
6,413,742 B1 6,428,978 B1		Olsen et al. Olsen et al.	2003/013336		12/2003	
6,440,167 B2	8/2002	Shimizu	2003/023274			Lamberti et al.
6,443,964 B1		Ory et al.	2004/0034373 2004/005440			Schuldt-Hempe et al. Dubson et al.
6,447,551 B1 6,447,802 B2		Goldmann Sessions et al.	2004/0059350			Gingras
6,448,378 B2	9/2002	DeVore et al.	2004/0101540	6 A1	5/2004	Gorman et al.
6,451,032 B1		Ory et al.	2004/0172043 2004/0215219			Browning Eldridge et al.
6,451,301 B1 6,454,787 B1		Sessions et al. Maddalo et al.	2005/0002893			Goldmann
6,477,865 B1		Matsumoto	2005/0010300	6 A1	1/2005	Priewe et al.
6,479,072 B1	11/2002	Morgan et al.	2005/0021058 2005/008592		1/2005	Negro Darois et al.
6,500,464 B2 6,509,031 B1		Ceres et al. Miller et al.	2005/008392			Popadiuk et al.
6,511,958 B1		Atkinson et al.	2005/011393	8 A1	5/2005	Jamiolkowski et al.
6,514,286 B1	2/2003	Leatherbury et al.	2005/0137512			Campbell et al.
6,514,514 B1 6,540,773 B2	2/2003 4/2003	Atkinson et al.	2005/014216 2005/0148963			Freeman et al. Brennan
6,541,023 B1		Andre et al.	2005/0175659		8/2005	Macomber et al.
6,548,077 B1	4/2003	Gunasekaran	2005/0228403			Fricke et al.
6,554,855 B1	4/2003	Dong Burgess et al.	2005/0232979 2005/024445:			Shoshan Greenawalt
6,559,119 B1 6,566,345 B2		Miller et al.	2005/026752	1 A1	12/2005	Forsberg
6,576,019 B1	6/2003	Atala	2005/028869		12/2005	Leiboff Matsuda et al.
6,596,304 B1 6,599,323 B2		Bayon et al. Melican et al.	2006/0094313 2006/013592			Wiercinski et al.
6,599,524 B2		Li et al.	2006/014750	1 A1	7/2006	Hillas et al.
6,599,690 B1		Abraham et al.	2006/016756			Odar et al.
6,613,348 B1 6,623,963 B1	9/2003	Jain Müller et al.	2006/0216320 2006/025298			Kitazono et al. Matsuda et al.
6,630,414 B1		Matsumoto	2007/0031474		2/2007	Tayot
6,637,437 B1*	10/2003	Hungerford et al 128/898	2007/0161109			Archibald et al.
6,638,284 B1 6,652,594 B2		Rousseau et al. Francis et al.	2007/0198040 2007/0280990		8/2007 12/2007	Buevich et al 606/151
6,653,450 B1		Berg et al.	2007/0280990	JAI	12/2007	Stopek
6,660,280 B1	12/2003	Allard et al.	F	OREIG	N PATE	NT DOCUMENTS
6,682,760 B2 6,706,684 B1		Noff et al. Bayon et al.				
6,706,690 B2		Reich et al.	DE DE	197 18 198 32		12/1997 1/1998
6,719,795 B1		Cornwall et al.	DE	197 51		12/1998
6,723,335 B1 6,730,299 B1		Moehlenbruck et al. Tayot et al.	DE	100 19	604	10/2001
6,743,435 B2		DeVore et al.	DE EP	100 43 0 248		6/2002 12/1987
6,752,834 B2*		Geistlich et al 623/23.63	EP	0 243		4/1988
6,773,723 B1 6,790,454 B1		Spiro et al. Abdul Malak et al.	EP	0 276		8/1988
6,833,408 B2		Sehl et al.	EP EP	0 194 0 372		11/1989 6/1990
6,835,336 B2	12/2004		EP	0 531		3/1993
6,852,330 B2 6,869,938 B1		Bowman et al. Schwartz et al.	EP	0 544	485	6/1993
6,893,653 B2		Abraham et al.	EP EP	0 552 0 611		7/1993 8/1994
6,896,904 B2	5/2005	Spiro et al.	EP	0 614		9/1994 9/1994
6,936,276 B2 6,939,562 B2		Spiro et al. Spiro et al.	EP	0 621	014	10/1994
6,939,362 B2 6,949,625 B2	9/2005	Tayot	EP EP	0 625 0 693		11/1994 1/1996
6,966,918 B1	11/2005	Schuldt-Hempe	EP	0 705		4/1996
6,974,679 B2		Andre et al.	EP	0 719	527	7/1996
6,974,862 B2	12/2005	Ringeisen et al.	EP	0 797	962	3/1997

(56)	Refer	ences Cited	WO WO 2004/078120 9/2004 WO WO 2005/013863 2/2005
	FOREIGN PAT	ENT DOCUMENTS	WO WO 2005/018698 3/2005
			WO WO 2005/105172 11/2005
$\mathbf{EP}$	0 774 240	5/1997	WO WO 2005/011280 12/2005
$\mathbf{EP}$	0 800 791	10/1997	WO WO 2006/018552 2/2006
EP	0 827 724	3/1998	WO WO 2006/023444 3/2006 WO WO 2007/048099 4/2007
EP	0 847 727	6/1998	WO WO 2007/048099 4/2007
EP EP	0 876 808	11/1998 2/1999	OTHER PUBLICATIONS
EP	0 895 762 0 898 944	3/1999	
EP	0 637 452	10/1999	Malette et al., Chitosan, a new hemostatic, Ann Th. Surg. 1983,
EP	1 017 415	7/2000	36:55-58.
EP	1 036 545	9/2000	Langenbech MR, et al., "Comparison of biomaterials in the early
EP	1 052 319	11/2000	postoperative period", Surg Endosc. 2003; 17(7):1105-9.
EP	1 055 757	11/2000	Bracco P., et al., "Comparison of polypropylene and polyethylene
EP EP	1 090 590 1 216 717	5/2001 6/2002	terephthalate (Dacron) meshes for abdominal wall hernia repair: a
EP	1 216 717	6/2002	chemical and morphological study", <i>Hernia</i> 2005, 9(1):51-55.
EP	0 836 838	6/2003	Klinge U, et al., "Foreign body reaction to meshes used for the repair
EP	1 315 468	6/2003	of abdominal wall hernias", Eur J. Surg 1999; 165:665-73.
$\mathbf{EP}$	1 382 728	7/2003	Logeart D, et al., "Fucans, sulfated polysaccharides extracted from brown seaweeds, inhibit vascular smooth muscle cell proliferation.
EP	1 484 070	12/2004	II. Degradation and molecular weight effect," Eur J Cell Biol. Dec.
EP EP	1 561 480	8/2005	1997; 74(4):385-90.
FR	1 782 848 2 244 853	5/2007 4/1975	Haneji K, et al., "Fucoidan extracted from Cladosiphon Okamuranus
FR	2 257 262	8/1975	Tokida induces apoptosis of human T-cell leukemia virus type 1-in-
FR	2 308 349	11/1976	flected T-cell lines and primary adult T-call leukemia cells", Nutrition
FR	2 453 231	10/1980	and Cancer, 2005; 52(2):189-201.
FR	2 612 392	9/1988	Junge K, et al, "Functional and morphologic properties of a modified
FR	2 715 405	7/1995	mesh for inguinal hernia repair", World J Surg 2002; 26:1472-80.
FR FR	2 724 563 2 730 406	3/1996 8/1996	Klinge, et al., "Functional and Morphological Evaluation of a Low-
FR	2 744 906	8/1997	Weight, Monofilament Polypropylene Mesh for Hernia Repair", J.
FR	2 766 698	2/1999	Biomed Mater Res 2002 63:129-136.
FR	2 771 622	6/1999	Welty G, et al., "Functional impairment and complaints following
FR	2 773 057	7/1999	incisional hernia repair with different polypropylene meshes", Her-
FR	2 774 277	8/1999	nia 2001;5:142-7.
FR FR	2 779 937 2 859 624	12/1999 3/2005	Varum KM et al., "In vitro degradation rates of partially N-acetylated
FR	2 863 277	6/2005	chitosans in human serum", Carbohydrate Research, 1997, 299:99-101.
FR	2 876 020	4/2006	Haroun-Bouhedja F, et al., "In vitro effects of furans on MDA-
FR	2 884 706	10/2006	MB231 tumor cell adhesion and invasion", Anticancer Res. JulAug.
GB	1 174 814	12/1969	2002;22(4):2285-92.
GB GB	2 051 153	1/1981	Scheidbach H, et al., "In vivo studies comparing the biocompatibility
JР	2 306 110 03032677	4/1997 2/1991	of various polypropylene meshes and their handling properties dur-
ĴР	05237128	9/1993	ing endoscopic total extraperitoneal (TEP) patchplast: an experimen-
JP	09137380	5/1997	tal study in pigs", Surg Endosc 2004; 18(2):211-20.
WO	WO 89/02445	3/1989	Blondin C, et al., "Inhibition of complement activation by natural
WO	WO 89/08467	9/1989	sulfated polysaccharides (fucans) from brown seaweed", Molecular
WO WO	WO 90/12551 WO 92/06639	11/1990 4/1992	Immuol. Mar. 1994; 31(4):247-53.
WO	WO 92/00039 WO 92/20349	11/1992	Zvyagintseva TN, et al., "Inhibition of complement activation by
wo	WO 93/11805	6/1993	water-soluble polysaccharides of some far-eastern brown seaweeds",
WO	WO 93/18174	9/1993	Comparative Biochem and Physiol Jul. 2000; 126(3):209-15. Rosen M, et al., "Laparoscopic component separation in the single-
WO	WO 94/17747	8/1994	stage treatment of infected abdominal wall prosthetic removal", Her-
WO WO	WO 95/07666 WO 95/18638	3/1995 7/1995	nia, 2007 11:435-440.
WO	WO 95/32687	12/1995	Amid P.K., "Lichtenstein tension-free hemioplasty: Its inception,
wo	WO 96/03091	2/1996	evolution, and principles", Hernia 2004; 8:1-7.
WO	WO 96/08277	3/1996	Boisson-Vidal C, et al., "Neoangiogenesis induced by progenitor
WO	WO 96/09795	4/1996	endothelial cells: effect of fucoidan from marine algae", Cardiovas-
WO	WO 96/14805	5/1996	cular & Hematological Agents in Medicinal Chem. Jan. 2007;
WO WO	WO 96/41588 WO 97/35533	12/1996 10/1997	5(1):67-77.
WO	WO 98/35632	8/1998	O'Dwyer PJ, et al., :Randomized clinical trial assessing impact of a
wo	WO 98/49967	11/1998	lightweight or heavyweight mesh on chronic pain after inguinal
WO	WO 99/05990	2/1999	hernia repair, Br J Surg. 2005; 92(2):166-70.
WO	WO 99/06080	2/1999	Muzzarelli, et al., "Reconstruction of parodontal tissue with
WO	WO 9906079	2/1999	chitosan", <i>Biomaterials</i> 1989, 10:598-604. Haroun-Bouhedja F, et al., "Relationship between sulfate groups and
WO	WO 99/51163	10/1999	biological activities of fucans", Thrombosis Res. Dec. 1, 2000,
WO WO	WO 00/16821 WO 01/15625	3/2000 3/2001	100(5):453-9.
WO	WO 01/13023 WO 01/80773	11/2001	Blondin C, et al., "Relationships between chemical characteristics
WO	WO 01/807/3 WO 02/07648	1/2002	and anticomplementary activity of fucans", <i>Biomaterials</i> . Mar. 1996;
wo	WO 02/078568	10/2002	17(6):597-603.
WO	WO 03/002168	1/2003	Strand SP, et al., "Screening of chitosans and conditions for bacterial
WO	WO 2004/004600	1/2004	flocculation", Biomacromolecules, 2001, 2:126-133.

#### (56)**References Cited**

# OTHER PUBLICATIONS

Kanabar V, et al., "Some structural determinants of the antiproliferative effect of heparin-like molecules on human airway smooth

muscle", *Br J Pharmacol*. Oct. 2005; 146(3):307-7. Hirano et al., "The blood biocompatibility of chitosan and N-acylchitosans", *J Biomed. Mater. Res.* 1985, 19:413-417. Rao S.B., et al., "Use of chitosan as a biomaterial: studies on its safety

and haemostatic potential", J. Biomed. Mater. Res. 1997, 34:21-28.

Prokop A, et al., "Water soluble polymers for immunoisolation I: complex coacevation and cytotoxicity", Advances in Polymer Science, 1998 136:1-51.

Collins et al., "Use of collagen film as a dural substitute: Preliminary Animal Studies", Journal of Biomedical Materials Research, vol. 25,

Preliminary Search Report from French Patent Office dated Dec. 20,

\* cited by examiner

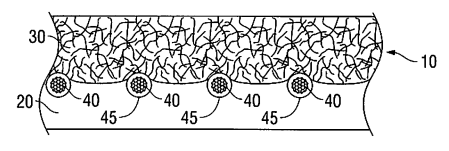


FIG. 1

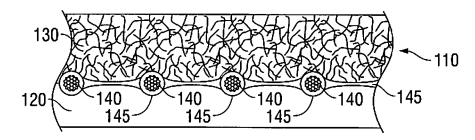
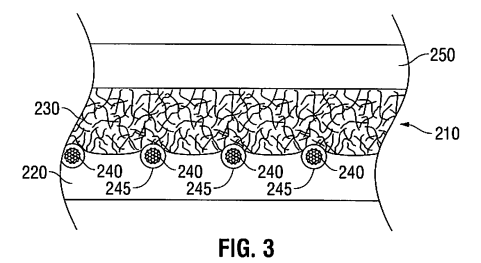


FIG. 2



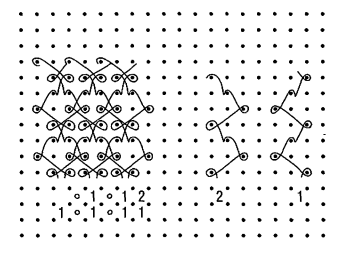
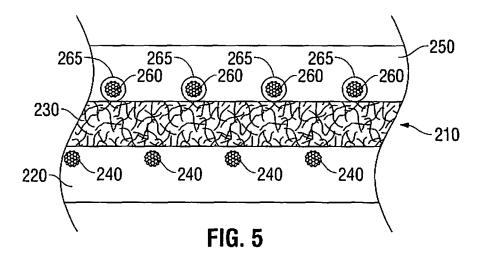
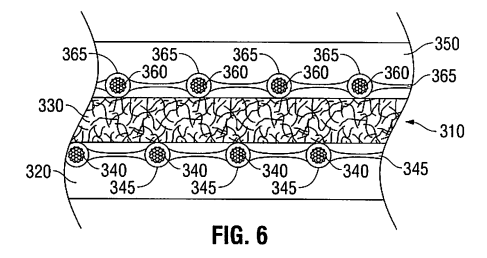


FIG. 4





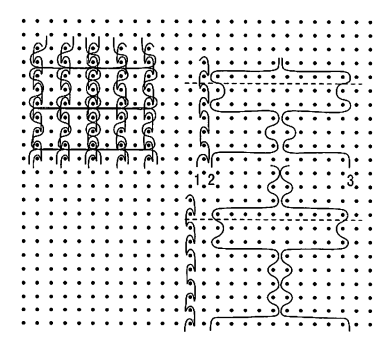


FIG. 7

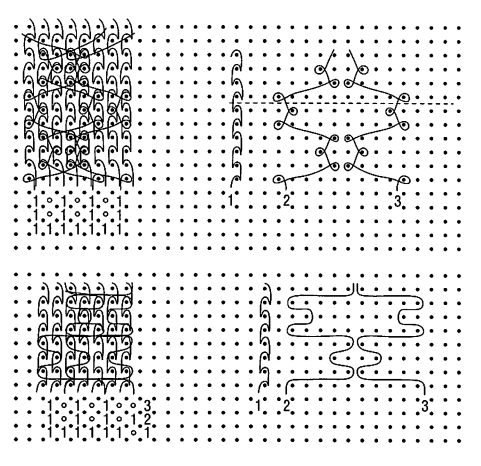


FIG. 8

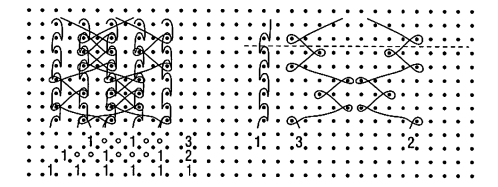


FIG. 9

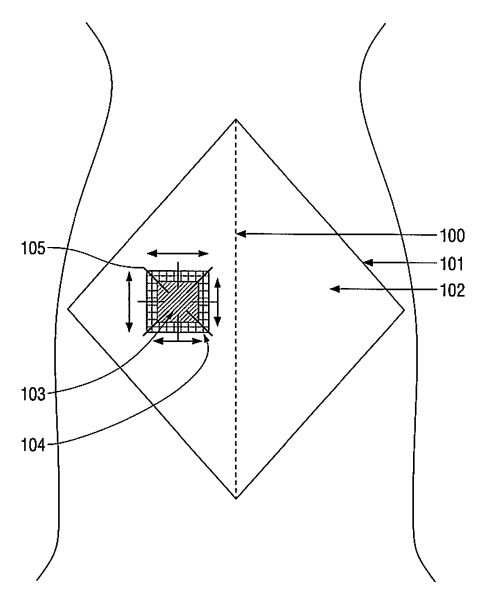


FIG. 10

# BIOSYNTHETIC IMPLANT FOR SOFT TISSUE REPAIR

# CROSS-REFERENCE TO RELATED APPLICATION

The present application claims priority to, and the benefit of, U.S. Provisional Patent Application Ser. Nos. 61/076,164 filed on Jun. 27, 2008 and 61/076,166 filed on Jun. 27, 2008, the entire disclosures of which are incorporated by reference herein.

# TECHNICAL FIELD

The present composite materials or implants have at least, a non-porous layer, a porous layer and a reinforcement member. The present composite materials or implants resist tearing when used in surgery, can simultaneously achieve hemostasis and prevent post-surgical adhesion wherever it is needed. They are particularly indicated for complex hernia, requiring resistance to microbial infection and a low speed of degradation during implantation. The present composite materials resist tearing when used in orthopedic surgery for tendon or ligament repair. They can support the repair/regeneration of soft tissues, under high stress/load.

#### DESCRIPTION OF THE RELATED ART

Implants for use in visceral surgery having a porous adhesive collagen layer closely associated with a collagen film are 3 known. In this type of material, the film helps prevent the formation of post-operative adhesions and the porous adhesive collagen layer functions as a hemostatic compress.

Such implants are frequently secured to tissue during surgery using a surgical fastener, such as a staple, clip, tack, suture or the like. Collagen, however, weakens quickly when exposed to the moist conditions within the body during surgery. As a result, previous composite implants are prone to tearing during implantation.

In addition, the strength of such implants typically is too 40 low for use under the high stresses or high loads encountered in orthopedic surgery, such as in tendon or ligament repair.

It would be advantageous to provide an implant having anti-adhesion, antimicrobial and relatively high strength properties and which resists tearing when subjected to the 45 forces associated with securing the implant to tissue using surgical fasteners.

It would be advantageous to provide an implant having anti-adhesion, hemostatic and antimicrobial properties and which resists tearing when subjected to the forces associated 50 with securing the implant to tissue using surgical fasteners.

In particular, it would be advantageous to provide an implant which maintains its properties, such as integrity, strength, resistance to tearing, porosity, thickness, when it is submitted to moist or wetted conditions, such as conditions 55 encountered after implantation, when the implant is surrounded by biological fluids. It would also be advantageous to provide an implant which, although bioabsorbable in the long term (i.e., over one year), is long lasting, i.e., maintains its properties and integrity for at least 6 months.

#### **SUMMARY**

The present implants therefore aim to considerably improve the previously described composite collagenic mate- 65 rials with respect to their handling characteristics and resistance to tearing during implantation, their time of full degra-

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dation (over 1 year), their ability to withstand high loads and their resistance to microbial contamination during implantation, in particular when they are submitted to moist conditions such as implantation conditions, when they are surrounded by biological fluids. These aims are achieved by the present implants which include at least a non-porous layer, a porous layer and a reinforcement member. In embodiments, the nonporous layer is an oxidized collagen constituent-containing film possessing anti-adhesion properties. In embodiments, the porous layer is a collagen-containing foam that provides a matrix for soft tissue repair and/or regeneration, and that is long lasting and maintains its properties and integrity even in a wetted state. In embodiments, the reinforcement member is a mesh, such as a mesh having a weight density of below about 50 g/m<sup>2</sup>. In other embodiments, the reinforcement member is a mesh, such as a mesh having a weight density of greater than about 100 g/m<sup>2</sup>. The mesh, which can be knitted either from biodegradable yarns, non biodegradable yarns or any combination of thereof, may be fully coated with a collagen based formulation so as to limit the inflammation reaction and the infection risk associated with the yarns of the mesh. In other embodiments, the mesh is coated by any materials which can reduce the inflammation reaction and the infection risk.

Methods for producing the present implants are also described. In embodiments, the reinforcement member is fully coated with cross-linked collagens. Then a liquid solution based on oxidized collagen destined to form the non-porous layer is cast on a substrate. The coated reinforcement member is applied to the solution, in embodiments becoming completely embedded therein, for example, by pressing the reinforcement member into the solution or by the application of additional solution on top of the original volume of solution. Prior to complete gelling, a pre-formed porous layer, made from long lasting collagenic formulation is laid on the surface of the gelling solution. Upon drying, the various components adhere to form the present implant.

One aspect of the present disclosure is an implant comprising:

- a porous layer comprising at least a collagen,
  - a non-porous layer comprising a collagenic constituent, the non-porous layer being joined to the porous layer, and a reinforcement member,
  - wherein the reinforcement member is embedded into the non-porous layer, and
  - the porous layer has a three dimensional density ranging from about 20 mg collagen/cm³ to about 200 mg collagen/cm³.

According to the present application, the three dimensional density in collagen of the porous layer is measured as follows: the amount of collagen used for the preparation of the porous layer is initially weighed in mg and recorded. The dimensions, i.e., length, width and thickness, of the porous layer in the final product, either in a dry or wetted state, are measured in cm. As meant in the present application, a wetted state corresponds to the soaking of the porous layer, or of the implant, during 24 hours in water at ambient temperature (i.e., 18-25° C.). The three dimensional density of the porous layer in mg collagen/cm³ results from the ratio of the amount weighed in mg over the volume measured in cm³.

The porous layer of the implant of the present disclosure has a three dimensional density ranging from about 20 mg collagen/cm³ to about 200 mg collagen/cm³, either in a dry or wetted state. As a consequence, the porous layer of the implant of the present disclosure has a three dimensional density ranging from about 20 mg collagen/cm³ to about 200 mg collagen/cm³, in a dry state. Also, the porous layer of the

implant of the present disclosure has a three dimensional density ranging from about 20 mg collagen/cm<sup>3</sup> to about 200 mg collagen/cm<sup>3</sup> in a wetted state, i.e., after soaking in water at ambient temperature (18-25° C.) during 24 hours. The porous layer of the implant of the present disclosure is therefore dense in collagen, i.e., it comprises a high ratio of collagen. Such a density allows the porous layer to be long lasting although the porous layer will degrade in vivo in the long term, i.e., over one year for example. For example, the porous layer of the implant of the present disclosure will not 10 be degraded before 6 months after the implantation. At the same time, the porous layer of the implant of the present disclosure shows a good homogeneity and favors the cellular growth. In particular, the porous layer of the implant of the present disclosure maintains its properties, such as integrity, 15 porosity, resistance to tearing, strength, thickness, even in moist conditions, i.e., in a wetted state, similar to the conditions of implantation, when the porous layer and/or the implant is surrounded by the biological fluids. In particular, the porous layer of the implant of the present disclosure 20 maintains substantially its thickness after implantation: for example there is no increase of 50% of the thickness or decrease of two thirds of the thickness of the porous layer after immersion of the porous layer in water, i.e., in moist conditions similar to the implantation moist conditions.

In an embodiment, the three dimensional density ranges from about 50 mg collagen/cm<sup>3</sup> to about 150 mg collagen/cm<sup>3</sup>.

In embodiments, the collagen of the porous layer comprises at least oxidized collagen.

In embodiments, the collagen of the porous layer further comprises a collagen crosslinked with a compound selected from glutaraldehyde, hexamethylene diisocyanate (HMDI) and mixtures thereof. For example, the compound is glutaraldehyde. Alternatively, the compound is hexamethylene diisocyanate.

In embodiments, the porous layer further comprises a chitosan.

In embodiments, the collagenic constituent of the non-porous layer is oxidized collagen.

In embodiments, the reinforcement member is a mesh having a weight density less than  $100 \, \mathrm{g/m^2}$ , for example less than  $50 \, \mathrm{g/m^2}$ . In such embodiments, in particular, the mesh may be isoelastic. By "isoelastic" is meant according to the present application that the mesh shows substantially similar elastic 45 properties in all directions. Such meshes are particularly suitable for implants of the present disclosure intended to be used as reinforcement implants for hernia repair for example.

In other embodiments, the reinforcement member is a mesh having a weight density greater than 50 g/m², for example greater than 100 g/m². In such embodiments, in particular, the mesh may be asymmetric. By "asymmetric" is meant in the present application that the mesh shows significantly different elastic properties in at least two different directions, for example in two perpendicular directions. Such meshes are particularly suitable for implants of the present disclosure intended to be used as reinforcement implants for tendon and/or ligament repair for example.

prises at least oxidize from glutaraldehyde, and mixtures thereof. In embodiments, the comprises a collagen from glutaraldehyde, and mixtures thereof. In embodiments, the comprises a chitosan. In embodiments, the comprises a chitosan.

In embodiments, the mesh may be coated with a collagen crosslinked with a compound selected from glutaraldehyde, 60 hexamethylene diisocyanate (HMDI) and mixtures thereof.

In other embodiments, the mesh may be coated with a mixture of oxidized collagen and chitosan.

In embodiments, the implant comprises an additional nonporous layer. The additional non-porous layer may have the 65 same composition or a different composition from the nonporous layer above. The additional non-porous layer may 4

comprise an additional reinforcement member embedded therein. The additional reinforcement member may be identical to or different from the reinforcement member above.

In embodiments, the porous layer is biodegradable. By biodegradable is meant herein that all the components forming the porous layer will be completely degraded in vivo after a certain amount of time. The porous layer of the implant of the present disclosure is long lasting and therefore will not be completely degraded in vivo before 6 months after implantation.

Another aspect of the present disclosure is a process for preparing an implant having a porous layer, a non-porous layer, and a reinforcement member comprising the steps of:

- a) Preparing a composition comprising at least a collagen, using purified fibers of collagen as starting material,
- b) Freeze drying the composition of a) to obtain a porous layer.
- c) Compressing the porous layer obtained in b) so as to obtain a three dimensional density ranging from about 20 mg collagen/cm³ to about 200 mg collagen/cm³,
- d) Pouring a solution of a collagenic constituent onto a support to obtain a non-porous layer,
- e) Applying a reinforcement member on a top surface of the non-porous layer obtained in d), and
- f) Applying the compressed porous layer obtained in c) on the top surface of the non-porous layer, during gelification of the non-porous layer.

The starting material used for obtaining the porous layer of
the process of the present disclosure is purified fibers of
collagen. Such purified fibers of collagen are obtained by
conventional techniques such as described in FR-A-2 715
309. For example, the collagen is extracted from porcine
dermis by solubilization at acidic pH or by digestion with
pepsin, and purified by saline precipitations according to
known techniques (see for example FR-A-2 715 309). Dry
collagen fibers are then obtained by precipitation of an acid
solution of such a collagen by adding NaCl, and then washing
and drying of the precipitate obtained with aqueous solutions
of acetone.

Using purified fibers of collagen for preparing the porous layer of the implant of the present disclosure allows determining exactly the content of collagen in the resulting porous layer of the final product.

In embodiments, the porous layer is compressed in step c) so as to obtain a three dimensional density ranging from about 50 mg collagen/cm³ to about 150 mg collagen/cm³.

In embodiments, the collagen of the porous layer comprises at least oxidized collagen.

In embodiments, the collagen of the porous layer further comprises a collagen crosslinked with a compound selected from glutaraldehyde, hexamethylene diisocyanate (HMDI) and mixtures thereof.

In embodiments, the collagen of the porous layer further comprises a chitosan.

In embodiments, the collagenic constituent of the nonporous layer is obtained from purified fibers of collagen used as starting material.

In embodiments, in step e), the reinforcement member is pressed into the solution of the non-porous layer, without causing significant disruption of the portion of the layer of solution in contact with the support, so as to embed the reinforcement member completely within the non-porous layer.

Alternatively, in step e), following application of the reinforcement member, additional solution of the collagenic constituent of step d) may be applied in an amount sufficient to

cover the reinforcement member, so as to embed the reinforcement member completely within the non-porous layer.

In embodiments, in step c), the porous layer is compressed so as to reduce its thickness from about 1.2 times to 5 times. For example, the porous layer obtained in c) has a thickness stranging from about 0.1 cm to 1.5 cm.

Another aspect of the present disclosure is an implant obtainable by a process as described above.

# BRIEF DESCRIPTION OF THE DRAWINGS

FIGS. 1-3 are schematic illustrations of composite materials in accordance with embodiments of the present disclosure:

FIG. 4 shows a knitting pattern suitable for making a mesh reinforcement member for use in embodiments described herein:

FIG. 5 is a schematic representation of a composite material in accordance with another embodiment of the present  $_{20}$  disclosure;

FIG. 6 is a schematic representation of a composite material in accordance with another embodiment of the present disclosure;

FIGS. **7-9** show illustrative knitting patterns suitable for 25 use in making reinforcement components that may be incorporated into composite materials in accordance with embodiments of the present disclosure; and

FIG. 10 is a schematic representation of an implanted implant in accordance with the present disclosure in a rat.

#### DETAILED DESCRIPTION

The present implants include one or more non-porous layers, a porous layer and one or more reinforcement members. As seen in FIG. 1, composite implant 10 includes non-porous layer 20, porous layer 30 and reinforcement members 40, which in this illustrative embodiment are multifilament yarns coated with coating 45 and embedded within non-porous layer 20. Each of these components and processes for preparing each component and the composite implant are described in greater detail below.

In the embodiment shown in FIG. 2, composite implant 110 includes non-porous layer 120, porous layer 130 and  $_{45}$  reinforcement members 140 which in this illustrative embodiment are multifilament yarns coated with coating 145 and embedded within non-porous layer 120.

In the embodiment shown in FIG. 3, composite implant 210 includes non-porous layers 220 and 250, porous layer 50 230 and reinforcement member 240 with coating 245, which in this illustrative embodiment is embedded within non-porous layers 220.

In the embodiment shown in FIG. 5, composite implant 210 includes non-porous layers 220 and 250, porous layer 55 230 and reinforcement members 240 and 260 coated with a coating 265, which in this illustrative embodiment are embedded within non-porous layers 220 and 250, respectively. In the embodiment shown in FIG. 6, composite implant 310 includes non-porous layers 320 and 350, porous layer 60 330 and reinforcement members 340 with coating 345, and 360 with coating 365, which in this illustrative embodiment are embedded within non-porous layers 320 and 350, respectively.

Each of these components and processes for preparing 65 each component and the composite implant are described in greater detail below.

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The Non-Porous Layer

The one or more non-porous layers may retard or prevent tissue ingrowth from surrounding tissues thereby acting as an adhesion barrier and preventing the formation of unwanted scar tissue. Thus, in embodiments, the one or more non-porous layers possesses anti-adhesion properties. By "non-porous layer" is meant according to the present application a layer having a surface substantially closed to all potential cellular growth, such as a film.

The one or more non-porous layers may be as well a physical barrier against microbial contamination.

The one or more (for example additional) non-porous layers of the present implant may be made from any biocompatible natural or synthetic material. The material from which each non-porous layer is formed may be the same or different and may be bioabsorbable or non-bioabsorbable. It should of course be understood that any combination of natural, synthetic, bioabsorbable and non-bioabsorbable materials may be used to form the one or more non-porous layers. Techniques for forming non-porous layers from such materials are within the purview of those skilled in the art and include, for example, casting, molding and the like.

Some non-limiting examples of materials from which a non-porous layer may be made include but are not limited to poly(lactic acid), poly(glycolic acid), poly(hydroxybutyrate), poly(phosphazine), polyesters, polyethylene glycols, polyethylene oxides, polyacrylamides, polyhydroxyethylmethylacrylate, polyvinylpyrrolidone, polyvinyl alcohols, polyacrylic acid, polyacetate, polycaprolactone, polypropylene, aliphatic polyesters, glycerols, poly(amino acids), copoly (ether-esters), polyalkylene oxalates, polyamides, poly(iminocarbonates), polyalkylene oxalates, polyoxaesters, polyorthoesters, polyphosphazenes and copolymers, block copolymers, homopolymers, blends and combinations thereof.

In embodiments, natural biological polymers are used in forming the one or more non-porous layers of the implant. Suitable natural biological polymers include, but are not limited to, collagen, gelatin, fibrin, fibrinogen, elastin, keratin, albumin, hydroxyethyl cellulose, cellulose, oxidized cellulose, hydroxypropyl cellulose, carboxyethyl cellulose, carboxymethyl cellulose, and combinations thereof. In addition, the natural biological polymers may be combined with any of the other polymeric materials described herein to produce a non-porous layer of the implant.

In embodiments, the non-porous layer of the implant comprises at least a collagenic constituent.

In such embodiments, an aqueous solution of a collagenic constituent may be used to form a non-porous layer of the present implants. As used herein, the term "collagenic constituent" designates collagen which has at least partially lost its helical structure through heating or any other method, or gelatine. The term "gelatine" here includes commercial gelatine made of collagen which has been denatured by heating and in which the chains are at least partially hydrolyzed (molecular weight lower than 100 kDa). The collagenic constituent used may advantageously be formed of non-hydrolyzed collagen, mainly composed of  $\alpha$  chains (molecular weight around 100 kDa). In the context of the present disclosure,  $\alpha$  chains means complete  $\alpha$  chains or fragments of these complete  $\alpha$  chains produced by the loss of a small number of amino acids. The term "non-hydrolyzed" as used herein means that less than 10% of the collagenic chains have a molecular weight below about 100 kDa. If heating is used to denature the helical structure of the collagen, the heating should be moderate and provided under gentle conditions so

as to avoid degradation by hydrolytic cleavage of the gelatine thus formed. Suitable gelatine materials are commercially available

The collagen used can be of human or animal origin. It may particularly be type I porcine or bovine collagen, or type I or 5 type III human collagen or mixtures in any proportions of the last two types. Native collagen may advantageously be used, in acid solution or after processing, to eliminate the telopeptides, notably by pepsin digestion. The collagen can also be oxidized: in such a case, the collagen may be modified by 10 oxidative cleavage using any technique know to those skilled in the art, including, but not limited to the use of periodic acid or one of its salts as described by Tardy et al. in U.S. Pat. No. 4,931,546, the entire content of which is herein incorporated by reference. Briefly, this technique involves mixing the col- 15 lagen in acid solution with a solution of periodic acid or one of its salts at a concentration of between about 1 and  $10^{-5}$  M, in embodiments between about 5 10<sup>-3</sup> and 10<sup>-1</sup> M, at a temperature of between about 10 and 25  $^{\circ}$  C. for about 10 minutes to 72 hours. This process breaks down hydroxylysine and the 20 sugars of the collagen, thus creating reactive sites without causing crosslinking. The oxidative cleavage of collagen allows moderate cross-linking later in the collagenic material. It should of course be understood that this function may be provided by other means of moderate cross-linking, for 25 example by beta or gamma irradiation, or other agents of moderate cross-linking, for example chemical reagents at suitably low and non-toxic doses.

In embodiments, at least one of the non-porous layers of the present implant is made from collagen modified by oxi-30 dative cleavage as described above. In other embodiments, the extent of collagen cross-linking can be increased by any techniques known to those skilled in the art to adjust the degradation time of the non-porous layer as desired. As used herein, the term "moderately crosslinked" means that the 35 degradation of the non-porous layer will be at least 90% complete (as measured by residual weight) by the end of about three weeks of implantation; the term "highly crosslinked" means that the degradation of the non-porous layer will be at least 90% complete (as measured by residual 40 weight) by the end of about three months of implantation; and the term "extremely highly crosslinked" means that the degradation of the non-porous layer will be at least 90% complete (as measured by residual weight) by the end of about two years of implantation.

In embodiments, the collagen is advantageously formed of non-hydrolyzed collagen, mainly composed of a chains (molecular weight around  $100\,\mathrm{kDa}$ ). In the context of the present disclosure,  $\alpha$  chains means complete  $\alpha$  chains or fragments of these complete  $\alpha$  chains produced by the loss of a small 50 number of amino acids. The term "non-hydrolyzed" as used herein means that less than 10% of the collagenic chains have a molecular weight below about  $100\,\mathrm{kDa}$ . If heating is used to denature the helical structure of the collagen, the heating should be moderate and provided under gentle conditions so 55 as to avoid degradation by hydrolytic cleavage of the gelatine thus formed.

In embodiments, the collagenic constituent is oxidized collagen.

In such embodiments, a solution of oxidized collagen as 60 defined above may be used to form the non-porous layer. Typically, a collagen concentration from about 5 g/l to about 50 g/l, in embodiments from about 25 g/l to about 35 g/l is used.

The solution of oxidized collagen may be heated, for 65 example to a temperature in excess of 37° C., in embodiments to a temperature of from about 40° C. to about 50° C., for at

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least one hour. This results in at least partial denaturing of the collagen's helical structure. Other physical or chemical techniques for denaturing collagen (e.g., ultrasonication, or by the addition of chaotropic agents) are within the purview of those skilled in the art may also be used.

In embodiments, the collagenic constituent of the nonporous layer is obtained from purified fibers of collagen used as starting material. Such purified fibers of collagen are obtained as described below for the porous layer.

In embodiments, at least one macromolecular hydrophilic additive that is chemically unreactive with the collagenic constituent may be added to the solution used to form the non-porous layer. "Chemically unreactive with the collagenic constituent" as used herein means a hydrophilic compound which is not likely to react with the collagenic constituent, notably which does not form covalent bonds with it during cross-linking.

The macromolecular hydrophilic additive advantageously has a molecular weight in excess of 3,000 Daltons, in embodiments from about 3,000 to about 20,000 Daltons. Illustrative examples of suitable macromolecular hydrophilic additives include polyalkylene glycols (such as polyethylene glycol), polysaccharides (e.g., starch, dextran and/or cellulose), oxidized polysaccharides, and mucopolysaccharides. It should of course be understood that combinations of macromolecular hydrophilic additives may be used. In embodiments, polyethyleneglycol 4000 (4000 corresponding to the molecular weight) is added as the macromolecular hydrophilic additive. The concentration of hydrophilic additive(s) can typically be from about 2 to about 10 times less than that of the collagenic constituent. Typically, the macromolecular hydrophilic additive is eliminated by diffusion through the non-porous layer, in a few days.

Optionally, glycerine may be added to the solution used to form the non-porous layer. When present, the concentration of glycerine in the solution can typically be from about 2 to about 10 times less than that of the collagenic constituent, in embodiments less than about one-third of the collagenic constituent concentration.

In illustrative embodiments of the solution used to form the non-porous layer, the concentrations of collagenic constituent, hydrophilic additive(s) and glycerine, when present, can be from about 2% to about 10% for the collagenic constituent, from about 0.6% to about 4% for the hydrophilic additive(s) and from about 0.3% to about 2.5% for glycerine, respectively.

The solution used to form the non-porous layer may be prepared by adding collagenic constituent, hydrophilic additive(s) and glycerine, when present, to water or a water/alcohol (e.g., ethanol) mixture at a temperature of from about 30° C. to about 50° C. The solution may advantageously be neutralized to a neutral pH to avoid hydrolyzing the collagenic constituent by heating and to obtain a film of physiological pH while permitting pre-cross-linking of the collagenic constituent if the mixture contains oxidized collagen as indicated previously.

The Porous Layer

The porous layer of the implant has a three dimensional density in collagen which may prevent or reduce the risk of microbial contamination of the implant. By "porous layer" is meant according to the present application a layer comprising pores, voids, holes allowing at least some cellular growth to take place.

The porous layer of the implant has for example openings or pores over at least a portion of a surface thereof. As described in more detail below, suitable materials for forming the porous layer include, but are not limited to foams (e.g.,

open or closed cell foams). In embodiments, the pores may be in sufficient number and size so as to interconnect across the entire thickness of the porous layer. In other embodiments, the pores do not interconnect across the entire thickness of the porous layer. Closed cell foams are illustrative examples of 5 structures in which the pores may not interconnect across the entire thickness of the porous layer. In yet other embodiments, the pores do not extend across the entire thickness of the porous layer, but rather are present at a portion of the surface thereof. In embodiments, the openings or pores are 10 located on a portion of the surface of the porous layer, with other portions of the porous layer having a non-porous texture. Those skilled in the art reading the present disclosure will envision other pore distribution patterns and configurations for the porous layer.

The porous layer of the present implant may include any biocompatible natural or synthetic material. The material contained in the porous layer may be bioabsorbable or nonbioabsorbable. It should of course be understood that any combination of natural, synthetic, bioabsorbable and non- 20 bioabsorbable materials may be used to form the porous layer. Some non-limiting examples of materials from which the porous layer may be made include but are not limited to poly(lactic acid), poly(glycolic acid), poly(hydroxybutyrate), poly(phosphazine), polyesters, polyethylene glycols, poly- 25 ethylene oxides, polyacrylamides, polyhydroxyethylmethylacrylate, polyvinylpyrrolidone, polyvinyl alcohols, polyacrylic acid, polyacetate, polycaprolactone, polypropylene, aliphatic polyesters, glycerols, poly(amino acids), copoly (ether-esters), polyalkylene oxalates, polyamides, poly(imi- 30 nocarbonates), polyalkylene oxalates, polyoxaesters, polyorthoesters, polyphosphazenes and copolymers, block copolymers, homopolymers, blends and combinations thereof.

In embodiments, the porous layer comprises one or more bioabsorbable, natural biological polymer. Suitable natural biological polymers include, but are not limited to, collagen, gelatin, cellulose, hydroxypropyl cellulose, carboxyethyl cellulose, chitin, chitosan, and combinations thereof. Alternatively, the polymer constituent may be a polysaccharide such as chitin or chitosan, or polysaccharides modified by oxidation of alcohol functions into carboxylic functions such as oxidized cellulose. In addition, the natural biological polymers may be combined with any biocompatible synthetic materials to produce the porous layer of the implant.

Where the porous layer is a foam, the porous layer may be formed using any method suitable to forming a foam or sponge including, but not limited to the lyophilization or freeze-drying of a composition. Suitable techniques for making foams are within the purview of those skilled in the art. 50

The porous layer can be at least about 0.1 cm thick, in embodiments from about 0.2 to about 1.5 cm thick. The porous layer can have a two dimensional density of not more than about 100 mg collagen/cm² and, in embodiments, from about 10 mg collagen/cm² to about 50 mg collagen/cm². The 55 two dimensional density of the porous layer is calculated by dividing the three dimensional density obtained as explained above by the thickness of the porous layer. The three dimensional density of the porous layer ranges from about 20 mg collagen/cm³ to about 200 mg collagen/cm³, in embodiments from about 50 mg collagen/cm³ to about 150 mg collagen/cm³, either in a dry state or a wetted state of the porous layer. The size of the pores in the porous layer can be from about 10 µm to about 500 µm, in embodiments from about 20 µm to about 200 µm.

Such a porous layer, in particular having a three dimensional density ranging from about 20 mg collagen/cm<sup>3</sup> to

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about 200 mg collagen/cm<sup>3</sup>, in embodiments from about 50 mg collagen/cm<sup>3</sup> to about 150 mg collagen/cm<sup>3</sup>, either in a dry or wetted state, is long lasting, i.e., it is not degraded before 6 months after implantation. At the same time, the porous layer of the implant of the present disclosure shows a good distribution and interconnection of the pores, thereby allowing an excellent cellular growth.

The porous layer of the implant of the present disclosure comprises at least a collagen. By "collagen" is meant according to the present application, a non-denatured collagen or collagen which has at least partially lost its helical structure through heating or any other method.

The starting material used for obtaining the porous layer of the process of the present disclosure is purified fibers of collagen. Such purified fibers of collagen are obtained by conventional techniques such as described in FR-A-2 715 309. For example, the collagen is extracted from porcine dermis by solubilization at acidic pH or by digestion with pepsin, and purified by saline precipitations according to known techniques (see for example FR-A-2 715 309). Dry collagen fibers are then obtained by precipitation of an acid solution of such a collagen by adding NaCl, and then washing and drying of the precipitate obtained with aqueous solutions of acetone.

In embodiments, the porous layer is a made from non-denatured collagen or collagen which has at least partially lost its helical structure through heating or any other method, consisting mainly of non-hydrolyzed  $\alpha$  chains, of molecular weight close to 100 kDa. The term "non-denatured collagen" means collagen which has not lost its helical structure. The collagen used for the porous layer of present implant may be native collagen or atelocollagen, notably as obtained through pepsin digestion and/or after moderate heating as defined previously. The origin and type of collagen may be as indicated for the non-porous layer described above.

In embodiments, the collagen of the porous layer comprises at least oxidized collagen. The oxidized collagen may be obtained as already described above for the non-porous layer.

In embodiments, the collagen may be cured to any desired degree. As used herein, the term "moderately cured" means that the degradation of the porous layer will be at least 90% complete (as measured by residual weight) by the end of about three weeks of implantation; the term "highly cured" means that the degradation of the porous layer will be at least 90% complete (as measured by residual weight) by the end of about three months of implantation; and the term "extremely highly cured" means that the degradation of the porous layer will be at least 90% complete (as measured by residual weight) by the end of about two years of implantation. In illustrative embodiments, moderately cured collagen is obtained by oxidative cleavage of collagen by periodic acid or one of its salts, as described for collagens of the non-porous layer. Highly cured collagen is made from collagen crosslinked by glutaraldehyde or by any other know cross-linking agents such as isocyanates, for example hexamethylene diisocyanate (HMDI). The degree of crosslinking distinguishes between highly cured and very highly cured materials. Techniques for curing to various degrees are within the purview of those skilled in the art.

In embodiments, the collagen of the porous layer further comprises a collagen crosslinked with a compound selected from glutaraldehyde, hexamethylene diisocyanate (HMDI) and mixtures thereof. For example, the compound is glutaraldehyde. Alternatively, the compound is hexamethylene diisocyanate.

In embodiments, the porous layer further comprises a chi-

The porous layer of the implant of the present disclosure may be obtained by freeze drying a composition comprising at least a collagen, using purified fibers of collagen as starting between the material.

In embodiments, the porous layer can be obtained by freeze-drying an aqueous acid solution or suspension of collagen at a concentration of about 2 to about 100 g/l and an initial temperature of about 4° C. to about 25° C. The concentration of collagen in the solution or suspension can be from about 10 g/l to about 100 g/l, in embodiments about 20 g/l.

The collagen suspension or solution can be made from 15 non-cured, moderately cured, highly cured or extremely highly cured collagens or combinations of thereof at any proportions. It may include as well non collagenic components, such as glycosaminoglycans, among them chitosan. The glycosaminoglycans display a degree of acetylation 20 (DA) of from about 0.5% to about 50%. They have a molecular weight ranging from about 100 to about 1,000,000 g/ml. It displays also a low polydispersity index of from about 1.2 to about 1.8. They may be a mixture of chitosans and other glycosaminoglycans (e.g. hyaluronic acid), which have free 25 amino groups capable of cross-linking to the oxidized collagen. In embodiments, the collagen suspension or solution is the combination of oxidized collagen and chitosan which can form a cross-linked network. As a result, in embodiments, the porous layer of the implant comprises oxidized collagen and chitosan, for example forming a network.

The porous layer is advantageously neutralized before its freeze-drying as a solution or suspension or after its freeze-drying under a dry form, at a pH from about 6 to about 8. In embodiments, the porous layer after its freeze-drying may be further cross-linked by any known cross-linking agents (e.g., glutaraldehyde, isocyanates) and/or by any physical treatment (e.g., thermal processing, gamma- and beta-irradiation).

The porous layer may be compressed so as to obtain a three dimensional density ranging from about 20 mg collagen/cm<sup>3</sup> to about 200 mg collagen/cm<sup>3</sup>, for example ranging from about 50 mg collagen/cm<sup>3</sup> to about 150 mg collagen/cm<sup>3</sup>.

For example, the porous layer may be further packed down by any relevant techniques so as to get a thickness from about 45 0.1 cm to about 1.5 cm and a 3D (three dimensional) density of from about 50 mg collagen/cm<sup>3</sup> to about 150 mg collagen/cm<sup>3</sup>. The porous layer may be compressed so as to reduce its thickness from about 1.2 times to 5 times.

In any case where the porous layer is packed down, it may be further stabilized by thermal treatment, in dry conditions, at a temperature up to about 100° C., for up to 2 days or by cross-linking with any known cross-linking agents. In other embodiments, it may be further stabilized by soaking the porous layer in a collagenic solution such as an oxidized collagen solution and by drying this solution. Agents in the collagen solution/suspension such as glycerin may be as well added, before the freeze-drying, to stabilize the thickness of the porous layer.

In embodiments, the porous layer is biodegradable. By biodegradable is meant herein that all the components forming the porous layer will be completely degraded in vivo after a certain amount of time. The porous layer of the implant of the present disclosure is long lasting and therefore will not be completely degraded in vivo before 6 months after implantation

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The Reinforcement Member

The present implant also includes one or more reinforcement members. The reinforcement member may be positioned entirely within the non-porous layer.

The reinforcement member can be a mesh prepared using any technique known to those skilled in the art, such as knitting, weaving, tatting, knipling or the like. Illustrative examples of suitable meshes include any of those that are presently commercially available for hernia repair. In embodiments where a mesh is used as the reinforcement member, the mesh will aid in affixing the composite to tissue without tearing of the porous or non-porous layers.

It is envisioned that the reinforcement member may be formed from any bioabsorbable, non-bioabsorbable, natural, and synthetic material previously described herein including derivatives, salts and combinations thereof. In particularly useful embodiments, the reinforcement member may be made from a non-bioabsorbable material to provide long term flexible tissue support. In embodiments, the reinforcement member is a surgical mesh made from polyester, polypropylene, poly(lactic acid) (PLA), poly(glycolic acid) (PGA), oxidized cellulose, polycaprolactone (PCL), polydiaxanone (PDO), trimethylene carbonate (TMC), polyvinyl alcohol (PVA), polyhydroxyalkanoates (PHAs), polyamides, polyethers, copolymers thereof and mixtures thereof. In addition polyethylene materials may also be incorporated into the implant described herein to add stiffness. Where monofilaments or multifilament braids are used as the reinforcement member, any commercially available suture material may advantageously be employed as the reinforcement member.

It is envisioned as well that the reinforcing member may be knitted from bioabsorbable yarns, non bioabsorbable yarns or both of them, in any combination.

In embodiments, the mesh is a flat mesh, essentially a two dimensional mesh, displaying a thickness lower than about 1 mm. The mesh may be isoelastic or asymmetric while displaying mechanical properties (stiffness, elasticity at physiological loads) close to the tissues to be repaired and/or regenerated.

In embodiments, the mesh weight density is below about 100 g per m<sup>2</sup>, in embodiments, below about 50 g per m<sup>2</sup>. The weight density of the mesh is determined according to the following method: the weight of the mesh is measured in grams; this weight is divided by the surface of the mesh in square meters. In such embodiments, the mesh may be isoelastic. Other characteristics which the reinforcement member may in embodiments advantageously possess include a breakage resistance of 100 N minimum (as determined by ISO 13934-1) and an extension breakage of 20 to 40% under 50 N (as determined by ISO 13934-1). Suitable fabrics suitable for obtaining isoelastic meshes can be made on a Rachel or warp knitting machine with at least two threading tools, one threaded full, one empty with PLA, PET or PP monofilament or multifilament biocompatible threads as shown in FIG. 4. When performed on a 22 or 24 gauge knitting machine, the macro-pores allow for a good integration to tissues. Such meshes are particularly suitable for implants of the present disclosure intended to repair an abdominal wall, for hernia treatment for example.

In other embodiments, the mesh is asymmetric, with a longitudinal axis displaying the highest mechanical properties and corresponding to the longitudinal axis of tendons and ligaments to be repaired.

In such embodiments, the ultimate load at break of the mesh in its longitudinal axis is from about 100 N to about 1,000 N, in embodiments over 250 N (as determined by ISO 13934-1).

In such embodiments, the elongation at break of the mesh in its longitudinal axis is less than about 100%, in embodiments less than about 20% (as determined by ISO 13934-1). The elongation at 100 N is typically from about 1% to about 20%, in embodiments from about 1% to about 10% (as determined by ISO 13934-1).

The pores size of the mesh are from about 1 mm to 5 mm, in embodiments from about 1 mm to 2.5 mm.

In embodiments, in particular when the mesh is asymmetric, the mesh weight density is greater than about  $50 \, g \, per \, m^2$ , 10 in embodiments over  $100 \, g \, per \, m^2$ .

In embodiments, the mesh is designed in such a way that it can be trimmed to the size of the defect to be repaired without any significant loss of the initial strain.

In embodiments, the mesh needs to hold sutures with a pull 15 out strength of at least 100 N (as determined by ISO 13934-1).

In embodiments, the porosity of the mesh is from about 20% to about 98%, in embodiments from about 40% to about 95%. For the purpose of the present application, the porosity of a mesh is intended to mean a porosity measured in the 20 following way: the dimensions, i.e., length, width and thickness, of the mesh, taken alone, are measured; moreover, the density of the yarns used to prepare the mesh is known. The mesh is weighed. By means of a simple subtraction, the volume occupied by the empty spaces within the mesh is 25 deduced therefrom. The porosity over the entire mesh is determined as being the percentage of empty volume relative to the total volume of the mesh.

In embodiments, a suitable two dimensional fabric for obtaining an asymmetric mesh is prepared on a Rachel or 30 warp knitting machine with three threading tool bars and a "marquisette" type weave, a bar forming a chain, two bars moving symmetrically as partial weft. (See FIG. 7) The number of crossings "under" a needle of these two weft bars determines the height of the opening. The three bars are one 35 threaded full, one empty 22 or 24 gauge with standard thread or high tenacity thread count mini 150 dtex monofilament or mini 50 dtex multifilament.

In other embodiments, another suitable two dimensional mesh is obtained with three knitting bars one of which is 40 threaded full, with the two design bars one full and one empty. (See FIG. 8.)

In yet other embodiments, another suitable two dimensional mesh is obtained with one full bar, one empty bar and two other one full and three empty (See FIG. 9).

Coating of the Reinforcement Member

In embodiments, the reinforcement member is coated, so as to cover at least some of the surfaces of the reinforcement member. In embodiments, most of the accessible surfaces of the reinforcement member are covered. In some embodiments, the resulting coated reinforcement member may appear partially or fully embedded in the coating formulation. It is envisioned that the reinforcement member may be coated with any bioabsorbable, non-bioabsorbable, natural, and synthetic material previously described.

In the embodiment shown in FIG. 2, composite implant 110 includes non-porous layers 120, porous layer 130 and reinforcement members 140 which in this illustrative embodiment are multifilament yarns coated with coating 145 and embedded within non-porous layer 120. Where two reinforcement members are used, for example, a reinforcement member and an additional reinforcement member, each reinforcement member may be coated and the coating compositions may be the same or different on each reinforcement member. It is also envisioned that where two reinforcement members are used, one may be coated and one may remain uncoated. In the embodiment shown in FIG. 5, for example,

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only reinforcement member 260 includes a coating 265 while reinforcement member 240 is uncoated. In the embodiment shown in FIG. 6, composite implant 310 includes non-porous layers 320 and 350, porous layer 330 and reinforcement members 340 with coating 345 and 360 with coating 365, which in this illustrative embodiment are embedded within non-porous layers 320 and 350, respectively.

In embodiments where the reinforcement member is a mesh, it may be coated with a collagenic coating by soaking the mesh once, twice, three times or more in a collagenic solution or suspension. The collagen can be native collagen or gelatin, as described above. The collagen also may be chemically modified, by oxidation, esterification (e.g., methylation, ethylation, succinylation) or the like. At the end of each soaking cycle, the collagen layer laid on the mesh may be further cross-linked by any known cross-linking agents such as glutaraldehyde and isocyanates. Alternatively, the collagen is cross-linked once at the end of the overall soaking process.

Where the reinforcement member is a mesh, the soaking collagen solution or suspension may include non collagenic components, such as glycosaminoglycans, among them chitosan, as described above in connection with the porous layer. The coating may be made from oxidized collagen and chitosan. The layer of the collagenic solution or suspension may be neutralized with an alkaline bath at the end of each soaking cycle or only once when the overall soaking process is completed.

In embodiments, the mesh is coated with a collagen crosslinked with a compound selected from glutaraldehyde, hexamethylene diisocyanate (HMDI) and mixtures thereof. In other embodiments, the mesh is coated with a mixture of oxidized collagen and chitosan.

In some embodiments, where the reinforcing member is a mesh, the mesh may be pretreated by plasma or gas techniques for the formation of a layer containing bioactive agents such as antimicrobial agents (e.g. DADMAC, Silver particles). For example, gas phase coating technology based on siloxane chemistry may be applied to the surface of the mesh where the siloxane thin film coating (<100 nm) material may include silver particles as described in the patents U.S. Pat. No. 6,984,392 and EP 1,691,606 B1.

Optional Bioactive Agents

In some embodiments, at least one bioactive agent may be combined with the implant and/or any of the individual components (the porous layer, the one or more non-porous layers, the one or more reinforcement members and/or the coating on the one or more reinforcement members) used to construct the implant. In these embodiments, the implant can also serve as a vehicle for delivery of the bioactive agent. The term "bioactive agent", as used herein, is used in its broadest sense and includes any substance or mixture of substances that have clinical use. Consequently, bioactive agents may or may not have pharmacological activity per se, e.g., a dye, or fragrance. Alternatively a bioactive agent could be any agent which provides a therapeutic or prophylactic effect, a compound that affects or participates in tissue growth, cell growth, cell differentiation, an anti-adhesive compound, a compound that may be able to invoke a biological action such as an immune response, or could play any other role in one or more biological processes. It is envisioned that the bioactive agent may be applied to the medical device in any suitable form of matter, e.g., films, powders, liquids, gels and the like.

Examples of classes of bioactive agents which may be utilized in accordance with the present disclosure include anti-adhesives, antimicrobials, analgesics, antipyretics, anesthetics, antiepileptics, antihistamines, anti-inflammatories, cardiovascular drugs, diagnostic agents, sympathomimetics,

cholinomimetics, antimuscarinics, antispasmodics, hormones, growth factors, muscle relaxants, adrenergic neuron blockers, antineoplastics, immunogenic agents, immunosuppressants, gastrointestinal drugs, diuretics, steroids, lipids, lipopolysaccharides, polysaccharides, and enzymes. It is also 5 intended that combinations of bioactive agents may be used.

Anti-adhesive agents can be used to prevent adhesions from forming between the implantable medical device and the surrounding tissues opposite the target tissue. In addition, anti-adhesive agents may be used to prevent adhesions from 10 forming between the coated implantable medical device and the packaging material. Some examples of these agents include, but are not limited to poly(vinyl pyrrolidone), carboxymethyl cellulose, hyaluronic acid, polyethylene oxide, poly vinyl alcohols and combinations thereof.

Antimicrobial agents may be included as a bioactive agent in the bioactive coating and/or in the film layers of the present disclosure to reinforce the antimicrobial properties of the implant of the present disclosure. Suitable antimicrobial agents include triclosan, also known as 2.4.4'-trichloro-2'- 20 hydroxydiphenyl ether, chlorhexidine and its salts, including chlorhexidine acetate, chlorhexidine gluconate, chlorhexidine hydrochloride, and chlorhexidine sulfate, biocide quaternary ammonium salts such as dimethyl diallyl ammonium chloride (DADMAC) and its derivatives, silver and its salts, 25 including silver acetate, silver benzoate, silver carbonate, silver citrate, silver iodate, silver iodide, silver lactate, silver laurate, silver nitrate, silver oxide, silver palmitate, silver protein, and silver sulfadiazine, polymyxin, tetracycline, aminoglycosides, such as tobramycin and gentamicin, 30 rifampicin, bacitracin, neomycin, chloramphenicol, miconazole, quinolones such as oxolinic acid, norfloxacin, nalidixic acid, pefloxacin, enoxacin and ciprofloxacin, penicillins such as oxacillin and pipracil, nonoxynol 9, fusidic acid, cephalosporins, and combinations thereof. In addition, antimicro- 35 bial proteins and peptides such as bovine lactoferrin and lactoferricin B and antimicrobial polysaccharides such as fucans and derivatives and oligomers of chitosan may be included as a bioactive agent in the bioactive coating of the present disclosure.

Other bioactive agents which may be included as a bioactive agent in the coating composition and/or in the film layer composition applied in accordance include: local anesthetics; non-steroidal antifertility agents; parasympathomimetic agents; psychotherapeutic agents; tranquilizers; deconges- 45 tants; sedative hypnotics; steroids; sulfonamides; sympathomimetic agents; vaccines; vitamins; antimalarials; anti-migraine agents; anti-parkinson agents such as L-dopa; antispasmodics; anticholinergic agents (e.g. oxybutynin); antitussives; bronchodilators; cardiovascular agents such as 50 coronary vasodilators and nitroglycerin; alkaloids; analgesics; narcotics such as codeine, dihydrocodeinone, meperidine, morphine and the like; non-narcotics such as salicylates, aspirin, acetaminophen, d-propoxyphene and the like; opioid receptor antagonists, such as naltrexone and naloxone; anticancer agents; anti-convulsants; anti-emetics; antihistamines; anti-inflammatory agents such as hormonal agents, hydrocortisone, prednisolone, prednisone, non-hormonal agents, allopurinol, indomethacin, phenylbutazone and the like; prostaglandins and cytotoxic drugs; estrogens; antibac- 60 terials; antibiotics; anti-fungals; anti-virals; anticoagulants; anticonvulsants; antidepressants; antihistamines; and immunological agents.

Other examples of suitable bioactive agents which may be included in the coating composition and/or in the film layer 65 composition include viruses and cells, peptides, polypeptides and proteins, analogs, muteins, and active fragments thereof,

such as immunoglobulins, antibodies, cytokines (e.g. lymphokines, monokines, chemokines), blood clotting factors, hemopoietic factors, interleukins (IL-2, IL-3, IL-4, IL-6), interferons ((3-IFN, (a-IFN and y-IFN), erythropoietin, nucleases, tumor necrosis factor, colony stimulating factors (e.g., GCSF, GM-CSF, MCSF), insulin, anti-tumor agents and tumor suppressors, blood proteins, gonadotropins (e.g., FSH, LH, CG, etc.), hormones and hormone analogs (e.g., growth hormone), vaccines (e.g., tumoral, bacterial and viral antigens); somatostatin; antigens; blood coagulation factors; growth factors (e.g., nerve growth factor, insulin-like growth factor); protein inhibitors, protein antagonists, and protein agonists; nucleic acids, such as antisense molecules, DNA and RNA; oligonucleotides; polynucleotides; and ribozymes. Assembling the Implant

In an illustrative embodiment, the implant is prepared by first pouring a solution of collagenic constituent, destined to form a non-porous layer, such as a film, possibly containing the hydrophilic additive(s) and glycerine, onto an adequate, substantially flat support and distributing it evenly.

The support is inert in that it does not react with the abovementioned components and is not involved in the cross-linking process. The support may advantageously be made from a hydrophobic material such as, for example, PVC or polystyrene. However, this support can also consist of a strippable material which will remain slightly adhesive and which can then be separated from the implant at the time of surgical use. This support may itself also consist of a film, for example dried collagen, onto which the solution is poured, or a layer of collagenic material gel in a distinctly more advanced state of gelification.

The density of the thin layer initially applied as a solution to the substrate can be from about 0.1 g solution/cm² to about 0.3 g solution/cm². This collagenic solution advantageously may be poured at a temperature from about 4° C. to about 30° C., and in embodiments from about 18° C. to about 25° C. Once applied to the substrate, the collagen solution is allowed to partially gel. Partial gelling results from cooling of the collagen solution, and not from drying of the solution.

A mesh reinforcement member is then applied to the solution. Application of the reinforcement member onto the solution means simply laying the reinforcement member onto the solution or partially gelled solution, and optionally applying slight pressing. The pressing should be insufficient to cause any significant disruption of the portion of the layer of solution in contact with the support/substrate thereby helping to maintain the integrity and anti-adhesion characteristics of the non-porous layer. The pressing may leave the surface of the reinforcement member exposed at the surface of the solution or may embed the reinforcement member completely within the layer of solution.

Following application of the mesh reinforcement member, but before complete gellification of the initially applied solution, additional solution may be applied in an amount sufficient to cover the mesh, so that it is completely embedded within the solution. Where pressing has already embedded the reinforcement member in the solution, application of additional solution may be eliminated.

This solution containing the embedded mesh reinforcement member is left to gel and a porous layer prepared as indicated above is applied to the solution during gelification.

Application of the porous layer onto the solution during gelification means simply laying the porous layer onto the gel, and optionally applying slight pressing. The pressing should be insufficient to cause any significant compaction of the porous layer. In embodiments where the porous layer has

been pre-formed, the porous layer will become joined to the solution, but will not become interlocked with the mesh reinforcement member.

The moment at which the porous layer is applied to the solution during gelification will depend upon the nature of the 5 solution employed, the conditions under which the solution is maintained during gelification and the nature of the porous layer. Generally, the solution will be allowed to gellify for a period of time prior to application of the porous layer such that the gel is still soft and allows the porous layer to penetrate 10 over a distance which is advantageously from about 0.05 mm to about 2 mm and, in embodiments from about around 0.1 mm to about 0.5 mm. The appropriate moment for application of the porous layer for any given combination of materials/ conditions can be determined empirically, for example by 15 applying small samples of the porous layer to the gel at various times and evaluating the degree of penetration and adherence. Generally, when the solution which is gelling is at a temperature of between about 4 and 30° C., the porous layer can be applied from about 5 to about 30 minutes after the 20 solution has been poured over the surface holding it.

The composite implant is left to dry or dried in order to obtain the final implant. When the collagenic solution destined to form the film includes oxidized collagen, it is polymerized while the material is drying. This drying occurs 25 favorably at a temperature of from about 4° C. to about 30° C., in embodiments from about 18° C. to about 25° C. In other embodiments, the implant is further processed by adding a second non-porous layer on the other side, as described above. The material can be dried in a jet of sterile air if 30 desired.

After drying, the implant can be separated from its support, packaged and sterilized using conventional techniques, e.g., irradiation with beta (electronic irradiation) or gamma (irradiation using radioactive cobalt) rays. In embodiments where 35 hydrolytically unstable materials are used in forming the composite, such as polyglycolic acid, polylactic acid the composites are packaged under sufficiently dry conditions to ensure that no degradation of the composite takes place during storage.

The present implants are stable at ambient temperature and remains stable for long enough to be handled at temperatures which may rise to temperatures of from about 37 to  $40^{\circ}$  C. The thickness of the non-porous layer is not critical, but typically can be less than about  $100~\mu m$  thick, and in embodiments 45 from about  $30~\mu m$  to about  $75~\mu m$  thick. Likewise, the thickness of the porous layer is not critical, but typically can be from about 0.1~cm to about 1.5~cm thick, and in embodiments from about 0.1~cm to about 1.2~cm thick. The implants in accordance with this disclosure can be produced at a desired 50~cm size or produced in large sheets and cut to sizes appropriate for the envisaged application.

The present composites may be implanted using open surgery or in a laparoscopic procedure. When implanted laparoscopically, the composite implant should be rolled with the 55 porous side on the inside before trocar insertion.

The porous layer of the present implant can act as a long lasting support of the repair and/or regeneration of any soft tissues. The high density of the porous layer—from about 20 mg/cm³ to about 200 mg/cm³—protects the healing wound to 60 some extent from bacteria and micro-organisms.

On the other hand, the implants described herein are particularly suitable for preventing post-operative adhesion, particularly in bleeding wounds, because the film prevents adherence. The non-porous layer also protects the healing wound for several days as it forms a barrier to bacteria and microorganisms.

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In embodiments where a mesh is used as a reinforcement member, the mesh will aid in affixing the composite to tissue without tearing of the porous or non-porous layers. The composite may be affixed to tissue using any conventional fastener, such as, for example, sutures, staples, tacks, two part fasteners, and the like. In embodiments, the fastener used to affix the composite to tissue is bioabsorbable, providing securement of the composite to a desired location long enough for tissue ingrowth to occur.

# **EXAMPLES**

The following non-limiting examples show possible combinations of the materials and their hemostatic powers and ability to prevent post-operative tissue adhesions.

#### Example 1

# 1°) Obtention of Purified Fibers of Collagen

The collagen used is porcine collagen type I, extracted from porcine dermis by solubilization at acidic pH or by digestion with pepsin, and purified by saline precipitations according to known techniques (see for example FR-A-2715 309). Dry collagen fibers obtained by precipitation of an acid solution of such a collagen by adding NaCl, and then washing and drying of the precipitate obtained with aqueous solutions of acetone having an increasing concentration of 80% to 100%, are used in the following steps of the present example.

# 2°) Preparation of Porous Layer

a) Preparation of Glutaraldehyde-Crosslinked Collagen

Purified fibers of collagen obtained in 1°) are solubilized in water at a final concentration of 1% m/v. The solution of collagen at 1% m/v is then neutralized by adding sodium phosphate at a final concentration of 20 mM. The final pH of the suspension was measured at between 6.5 and 7.5. Glutaraldehyde (aqueous solution of glutaraldehyde at 25%, m/v, sold by the company Fluka Chemie GmbH, Buchs, Switzerland) is then added to the suspension at a final concentration of 0.5% m/v. After two hours at ambient temperature, collagen fibers are recovered by filtration of the suspension through a nylon mesh. These fibers are then treated with sodium borohydride for at least two hours until the yellow coloration of the fibers has completely disappeared. The white fibers thus obtained are washed and neutralized at pH 6.5-7.5, and dried by removing the water with acetone. The acetone residues are then evaporated off.

# b) Preparation of Oxidized Collagen

A 30 g/l solution of collagen is prepared by dissolving purified fibers of collagen obtained in 1°) in 0.01 N HCl. Its volume is 49 liters. Periodic acid is added to it at a final concentration of 8 mM, i.e., 1.83 g/l. Oxidation takes place at an ambient temperature close to 22° C. for 3 hours away from light.

Then an equal volume of a solution of sodium chloride is added to the solution to obtain a final concentration of 41 g/l NaCl

After waiting for 30 minutes, the precipitate is collected by decantation through a fabric filter, with a porosity close to 100 microns, then washed 4 times with a 41 g/l solution of NaCl in 0.01 N HCl. This produces 19 kg of acid saline precipitate. This washing process eliminates all traces of periodic acid or iodine derivatives during oxidation of the collagen.

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Then, several washes in an aqueous solution of 80% acetone are used to concentrate the collagen precipitate and eliminate the salts present.

A final wash in 100% acetone is used to prepare 3.6 kg of a very dense acetone precipitate of acid, oxidized, non-reticulated collagen, with no trace of undesirable chemical products.

The acetone paste is diluted with apyrogenic distilled water at ambient temperature, to obtain a 2% concentration of collagen.

c) Preparation of a Composition of Collagen Under the Form of a Collagenic Suspension

A composition under the form of a suspension of collagen is prepared by mixing the glutaraldehyde-crosslinked collagen obtained in a) above and the oxidized collagen obtained in b) above, at the following concentrations:

- 0.5 to 3% m/v of glutaraldehyde-crosslinked collagen,
- 0.2 to 2% m/v of oxidized collagen.
- d) Preparation of the Porous Layer by Freeze-Drying

The collagen suspension thus obtained in c) above is then lyophilized according to the following method: freezing is carried out as rapidly as possible, by decreasing the temperature of the product from 8° C. to -45° C., generally in less than 2 hours. Primary desiccation is initiated at -45° C., at a 25 pressure of from 0.1 to 0.5 mbar. During this step, the temperature is gradually increased, with successive slopes and plateau, to +30° C. The lyophilization ends with secondary desiccation, at +30° C., for 1 to 24 hours. The vacuum at the end of secondary desiccation is between 0.005 and 0.2 mbar. 30 The total lyophilization time is from 18 to 72 hours.

After the freeze-drying, the porous layer is packed down, in other words compressed, so as to reduce its thickness from about 1.2 times to about 5 times and so as to give the porous layer a three dimensional density ranging from 20 mg collagen/cm³ to about 200 mg collagen/cm³ either in the dry or wetted state.

The porous layer may be optionally treated with a solution of hexamethylene diisocyanate (HMDI) 0.1% w/v in acetone. The treatment period is about 20 hrs. The porous layer is then 40 washed several times with acetone. The solvent is then eliminated by evaporation. HMDI may be replaced by any other suitable cross-linking agent such as glutaraldehyde, isocyanates (e.g. isophorone diisocyanate), bifunctional or trifunctional glycidyl ethers, carbodiimides, acyl azides, divinylsul-45 phone.

In any case, the porous layer is then heated at  $50^{\circ}$  C. for a period lasting between 15 and 24 hours to improve the cohesion and mechanical resistance of the lyophilized product during assembly of the composite.

# 3°) Preparation of a Solution of Oxidized Collagen Used to Form a Non-Porous Film

Oxidized collagen is prepared as described above in 2°)b) 55 in this example, but with minor modifications, at the end of the manufacturing process. The acetone paste is diluted with apyrogenic distilled water at 40° C., to obtain a 3% concentration of collagen, for a volume of 44 liters. The collagen suspension of a volume of 44 liters is heated for 30 minutes at 60 50° C., then filtered under sterile conditions through a membrane of 0.45 micron porosity in a drying oven at 40° C.

As soon as this solution is homogeneous and at 35° C., a sterile concentrated solution of PEG 4000 (polyethylene glycol with a molecular weight of 4000 Daltons) and glycerine is 65 added to it to produce a final concentration of 0.9% PEG, 0.54% glycerine and 2.7% oxidized collagen.

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As soon as these additions have been made, the pH of the solution is adjusted to 7.0 by adding a concentrated solution of sodium hydroxide.

#### 4°) Coating of the Reinforcement Member

A knitted isoelastic, multifilament polylactic acid/polyester mesh reinforcement member is coated in a solution of type I porcine collagen at 0.8 m/v, by soaking it in the solution, with a padder, machine usually employed in the textile industry for coating process. This cycle of processes is repeated up to three times in order to obtain the full covering of the mesh.

At the end of the coating, the collagen deposited on the knit is cross-linked with glutaraldehyde at 0.5% m/v (aqueous solution of glutaraldehyde at 25%, m/v, sold by the company Fluka Chemie GmbH, Buchs, Switzerland), at neutral pH (pH between 6.5 and 7.5), for 2 hours, and is then reduced with sodium borohydride. The reagents used are removed by washing the knit with several water baths.

The cross-linking of the collagen deposited on the knit can alternatively be carried out at the end of each coating cycle.

# 5°) Preparation of a Biosynthetic Mesh for Soft Tissue Repair

An implant having a porous layer made from a composition that includes a collagen joined to a fiber-reinforced film made from a composition that includes a collagenic constituent is prepared. The collagen solution destined to form the non-porous layer, as described in 3°) above, is poured in a thin layer on a framed, flat hydrophobic support such as PVC or polystyrene, at an ambient temperature close to 22° C. The amount of solution used is 0.133 grams of solution per square centimeter of support. The collagen layer is let for about 15 min under a laminar flow hood. Then, the collagen coated reinforcement member obtained in 4°) above is pressed in the gelled collagen layer. The pre-made porous layer obtained in 2°) is then applied and adheres to the still sticky surface of the collagen solution.

The composite material is then dehydrated in a drying cabinet at  $20^{\circ}$  C. and 40% humidity with a horizontal flow of filtered air at a velocity of  $1.2 \text{ m}^2/\text{s}$ .

An implant is obtained, which is long lasting and resists tearing when put under moist conditions similar to conditions encountered during implantation. In particular, the implant of the present example maintains its integrity, its strength and its resistance to tearing even after being it soaked in water during 24 hours at ambient temperature (18-25° C.). Such an implant may be used for hernia repair.

# Example 2

The product of Example 1 is further processed by adding a second non-porous layer on the other side. This second non-porous layer includes a second reinforcement member.

The collagen solution destined to form the second non-porous layer, as described in above Example 1, part 3°), is poured in a thin layer on a framed, flat hydrophobic support such as PVC or polystyrene, at an ambient temperature close to 22° C. The amount of solution used is 0.133 grams of solution per square centimeter of support. The collagen layer is let for about 15 min under a laminar flow hood. Then, a second collagen coated reinforcement member is pressed in the partially gelled collagen layer. Finally, the porous layer of the final product of Example 1 is laid on the partially gelled, second non-porous layer, in which the second reinforcement member is embedded.

The composite material is then dehydrated in a drying cabinet at  $20^{\circ}$  C. and 40% humidity with a horizontal flow of filtered air at a velocity of  $1.2 \text{ m}^2/\text{s}$ .

An implant is obtained, which is long lasting and resists tearing when put under moist conditions similar to conditions encountered during implantation. In particular, the implant of the present example maintains its integrity, its strength and its resistance to tearing even after being it soaked in water during 24 hours at ambient temperature (18-25° C.). Such an implant may be used for hernia repair.

#### Example 3

# 1°) Preparation of Porous Layer

A composition of a collagen/chitosan mixture was prepared by mixing an acidic solution of oxidized collagen and an acidic solution of chitosan in different proportions with a final polymer concentration of 1% (w/w).

#### a) Oxidized Collagen

Oxidized collagen was obtained as described in the porous layer description of the Example 1, part 2°)b). A 2% w/v solution of oxidized collagen was obtained from the collagen paste, at ambient temperature.

#### b) Chitosan

The chitosan was solubilized in deionized water with a stoichiometric amount of hydrochloric acid with a polymer concentration of 3% (w/w). The pH of the chitosan solution was about 5, but the pH could have been adjusted to 3 to have 30 better control of the crosslink kinetic between the oxidized collagen and chitosan.

#### c) Freeze Dried Composite

Several mixtures of various blends of oxidized collagen and chitosan as well as native collagen and chitosan (approximately 180 g) were obtained. Optionally, bioactive agents are included, such as antimicrobial agents. To the collagen preparation, may be added glycerol up to 1% w/v. The mixtures were poured within a 12 cm by 17 cm plastic box and freezedried for 48 hours. The samples were then neutralized in a 1M sodium hydroxide bath for 1 hour and thoroughly washed in deionized water until the pH reached 7. The freeze-dried sponges were then packed down, i.e., compressed, to obtain a porous layer with a final thickness of 3 mm and a three dimensional density ranging from 20 mg collagen/cm³ to 45 about 200 mg collagen/cm³, either in the dry or wetted state.

Additives, such as fucans, oligomers of chitosan, native or chemically modified glucosaminoglycans, which may induce self chemical crosslink between collagen and glucosaminoglycans (hyaluronic acid, sulphate chondroitin, etc.), oxidized starch, and any other product which may enhance tissue repair, limit the risk of sepsis, and modulate the mechanical properties of the composite (swelling rate in water, tensile strength, etc.) could have been added to the blend of oxidized collagen and chitosan.

The porous layer is then heated at a temperature equal or above 50° C. for a period lasting between 15 and 24 hours to improve the cohesion and mechanical resistance of the lyophilized product during assembly of the composite.

# 2°) Coating of the Reinforcement Member

A knitted isoelastic, multifilament polylactic acid/polyester mesh reinforcement member is soaked once, twice, or three times in an oxidized collagen/chitosan mixture with a 65 padder, then dried and neutralized with an alkaline bath so as to cover the accessible surface of the mesh yarns. Alterna-

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tively, layers of oxidized collagen/chitosan mixture may be neutralized after each soaking cycles.

# 3°) Preparation of a Biosynthetic Mesh for Soft Tissue Repair

An implant having a porous layer made from a composition that includes a cross-linked network of oxidized collagen and chitosan joined to a fiber-reinforced film made from a composition that includes a collagenic constituent is prepared. The collagen solution destined to form the non-porous layer, as described in the Example 1, part 3°), is poured in a thin layer on a framed, flat hydrophobic support such as PVC or polystyrene, at an ambient temperature close to 22° C. The amount of solution used is 0.133 grams of solution per square centimeter of support. The collagen layer is let for about 15 min under a laminar flow hood. Then, the collagen coated reinforcement member as obtained in 2°) above is pressed in the gelled collagen layer. The pre-made, compressed porous  $^{20}~$  layer as obtained in  $1^{\circ}) d)$  above is then applied and adheres to the still sticky collagen solution of the non-porous layer exposed through the pores of the mesh.

The composite material is then dehydrated in a drying cabinet at  $20^{\circ}$  C. and 40% humidity with a horizontal flow of filtered air at a velocity of  $1.2 \text{ m}^2/\text{s}$ .

The composite material may be optionally treated with a solution of hexamethylene diisocyanate (HMDI) 0.1% w/v in isopropanol. The treatment period is about 20 hrs. The porous layer is then washed several times with acetone. The solvent is then eliminated by evaporation.

HMDI may be replaced by any other suitable cross-linking agent such as glutaraldehyde, other isocyanates, bifunctional or trifunctional glycidyl ethers, carbodiimides, acyl azides, divinylsulphone.

The porous layer can be further heated at about 50° C. for a period lasting between 15 and 24 hours to improve the cohesion and mechanical resistance of the lyophilized product during assembly of the composite

An implant is obtained, which is long lasting and resists tearing when put under moist conditions similar to conditions encountered during implantation. In particular, the implant of the present example maintains its integrity, its strength and its resistance to tearing even after being it soaked in water during 24 hours at ambient temperature (18-25° C.). Such an implant may be used for hernia repair.

#### Example 4

# 1°) Collagen Used

The collagen used are purified fibers of collagen as described in example 1, part  $1^{\circ}$ ).

# 2°) Preparation of Porous Layer

a) Preparation of a Composition of Hexamethylene Diisocyanate (HMDI)-Crosslinked Collagen

Fifty grams of dry purified fibers of porcine collagen of 1°) above are mixed with 1 liter of acetone. One gram of HMDI is then added to the collagen suspension. The mixture is let, under agitation, overnight, at ambient temperature. Collagen fibres are then recovered by filtration of the suspension through a nylon mesh and are thoroughly washed with dry acetone to remove HMDI and acetone soluble HMDI byproducts. The cross-linked collagen fibres thus obtained are dried by removing the acetone residues. They may be further ground.

b) Preparation of Oxidized Collagen

Oxidized collagen is prepared from porcine collagen as described in Example 1, part 2°)b).

c) Preparation of the Collagenic Suspension

A composition under the form of a suspension of collagen 5 is prepared by mixing the HMDI-crosslinked collagen of a) above, the oxidized collagen of b) above and, optionally glycerol, at the following concentrations:

0.5 to 3% w/v of HMDI-crosslinked collagen,

0.2 to 2% w/v of oxidized collagen,

0 to 1% w/v of glycerol

d) Preparation of the Porous Layer by Freeze-Drying

Several compositions as above in c) under the form of mixtures of various blends of oxidized collagen and HMDI-cross-linked collagen (approximately 180 g) were poured within a 12 cm by 17 cm plastic box and freeze-dried for 48 hours.

After the freeze-drying, the porous layer is packed down, i.e., compressed, so as to reduce its thickness from about 1.2 times to about 5 times and so as to obtain a three dimensional density ranging from 20 mg collagen/cm³ to about 200 mg collagen/cm³, either in the dry or wetted state.

The porous layer may be optionally treated with a solution of hexamethylene diisocyanate (HMDI) 0.2% w/v in acetone. The treatment period is about 20 hrs. The porous layer is then 25 washed several times with acetone. The solvent is then eliminated by evaporation.

HMDI may be replaced by any other suitable cross-linking agent such as glutaraldehyde, isocyanates, bifunctional or trifunctional glycidyl ethers, carbodiimides, acyl azides, divinylsulphone.

In any case, the porous layer is then heated at a temperature above 50° C. for a period lasting between 15 and 24 hours to improve the cohesion and mechanical resistance of the lyophilized product during assembly of the composite.

#### 3°) Coating of the Reinforcement Member

A knitted isoelastic, multifilament polylactic acid/polyester mesh reinforcement member is coated in a solution of type 40 I porcine collagen at 0.8% m/v, by soaking it in the solution, with a padder, machine usually employed in the textile industry for coating process. This cycle of processes is repeated up to three times in order to obtain the full covering of the mesh.

At the end of the coating, the collagen deposited on the knit 45 is cross-linked with HMDI. The knit is crosslinked in 100 ml of a phosphate buffer (0.054 M Na<sub>2</sub>HPO<sub>4</sub>, 0.013 M NaH<sub>2</sub>PO<sub>4</sub>, pH 7.4) containing 3% (w/w) HMDI and 1.0% (w/w) Tween 80 (polyoxyethylenesorbitan monooleate) as a surfactant for 20 h at room temperature. The knits are then 50 rinsed for 30 min by running demineralized water, washed twice for 30 min with 4 M NaCl and washed four times for 30 min with distilled water to remove unreacted HMDI or surfactant before drying the knit.

The cross-linking of the collagen deposited on the knit can staternatively be carried out at the end of each coating cycle.

# 4°) Preparation of a Biosynthetic Mesh for Soft Tissue Repair

An implant having a foam layer (porous layer) made from a composition that includes a collagen joined to a fiber-reinforced film made from a composition that includes a collagenic constituent is prepared. The collagen solution destined to form the non-porous layer, as described in the 65 Example 1, part 3°) is poured in a thin layer on a framed, flat hydrophobic support such as PVC or polystyrene, at an ambi-

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ent temperature close to  $22^{\circ}$  C. The amount of solution used is 0.133 grams of solution per square centimeter of support. The collagen layer is let for about 15 min under a laminar flow hood. Then, the collagen coated reinforcement member of  $3^{\circ}$ ) above is pressed in the gelled collagen layer. The pre-made, compressed porous layer obtained in  $2^{\circ}$ ) above is then applied and adheres to the still sticky collagen solution exposed through the pores of the mesh.

The composite material is then dehydrated in a drying cabinet at  $20^{\circ}$  C. and 40% humidity with a horizontal flow of filtered air at a velocity of  $1.2 \text{ m}^2/\text{s}$ .

A second non-porous layer may be further prepared by pouring the collagen solution as above in a thin layer on a framed, flat hydrophobic support such as PVC or polystyrene, at an ambient temperature close to 22° C. The amount of solution used is 0.133 grams of solution per square centimeter of support. The collagen layer is let for about 15 min under a laminar flow hood. Then, the porous layer of the composite material obtained above is pressed in the gelled collagen layer.

The resulting material is then dehydrated in a drying cabinet at  $20^{\circ}$  C. and 40% humidity with a horizontal flow of filtered air at a velocity of  $1.2 \text{ m}^2/\text{s}$ .

An implant is obtained, which is long lasting and resists tearing when put under moist conditions similar to conditions encountered during implantation. In particular, the implant of the present example maintains its integrity, its strength and its resistance to tearing even after being it soaked in water during 24 hours at ambient temperature (18-25° C.). Such an implant may be used for hernia repair.

# Example 5

The product of Example 1 is further processed by adding a second non-porous layer on the other side.

The collagen solution destined to form the second non-porous layer, as described in Example 1, part  $3^{\circ}$ ), is poured in a thin layer on a framed, flat hydrophobic support such as PVC or polystyrene, at an ambient temperature close to  $22^{\circ}$  C. The amount of solution used is 0.133 grams of solution per square centimeter of support. The collagen layer is let for about 15 min under a laminar flow hood. Then, the porous layer of the final product of Example 1 is laid on the partially gelled second non-porous layer.

The composite material is then dehydrated in a drying cabinet at  $20^{\circ}$  C. and 40% humidity with a horizontal flow of filtered air at a velocity of  $1.2 \text{ m}^2/\text{s}$ .

An implant is obtained, which is long lasting and resists tearing when put under moist conditions similar to conditions encountered during implantation. In particular, the implant of the present example maintains its integrity, its strength and its resistance to tearing even after being it soaked in water during 24 hours at ambient temperature (18-25° C.). Such an implant may be used for hernia repair.

#### Example 6

# 1°) Collagen Used

The collagen used are purified fibers of collagen as described in example 1, part 1°).

# 2°) Preparation of the Porous Layer

 a) Preparation of a Composition of HMDI-Crosslinked Collagen

Fifty grams of dry fibers of porcine collagen as obtained in 5 1°) above are mixed with 5 liters of demineralized water, under agitation. Once the solution is clear, the collagen is further neutralized and precipitated with 1N sodium hydroxide solution, at a pH ranging from 6.5 to 7.5. The collagen precipitate is collected by centrifugation (10 min, 10,000 10 rpm). The precipitate is washed by 3 successive acetone baths and is then completely dried. The yield of the neutralizing step of collagen is generally over 90%.

Forty grams of dry neutralized collagen is mixed with 0.8 liter of acetone. Two thousand and four hundred milligrams of 15 HMDI is then added to the collagen suspension. The mixture is let, under agitation, from 6 to 20 hrs, at ambient temperature, under mild agitation. Collagen fibres are then recovered by filtration of the suspension through a nylon mesh and are thoroughly washed with dry acetone to remove HMDI and 20 acetone soluble HMDI byproducts. This HMDI cross-linking step can be repeated once or twice. At the end, the cross-linked collagen fibres thus obtained are dried by removing the acetone residues. If needed, they may be further milled.

A test confirmed that the obtained HMDI cross-linked 25 collagen is not substantially degraded by collagenase (Sigma), after 1 day in contact with collagenase, at +37° C. Generally, the degradation of the obtained HMDI cross-linked collagen by collagenase is less than 5% of its original amount whereas 65% or more of non cross-linked collagen is 30 degraded in the very same conditions of the collagenase test.

b) Preparation of Oxidized Collagen

Oxidized collagen is prepared from purified fibers of porcine collagen as described in the Example 1, part 2°)b).

c) Preparation of a Composition Under the Form of a 35 Collagenic Suspension

A suspension of collagen is prepared by mixing the HMDI-cross-linked collagen of a) above, and the oxidized collagen of b) above, at the following concentrations:

- 0.7 to 3% w/v of HMDI cross-linked collagen,
- 0.2 to 1.5% w/v of oxidized collagen.
- d) Preparation of the Porous Layer by Freeze-Drying

Several compositions as above in c), under the form of mixtures of various blends of oxidized collagen and HMDI-cross-linked collagen (approximately 180 g) were poured 45 within a 12 cm×17 cm plastic box and freeze-dried for 48 hrs.

After the freeze-drying, the porous layer is further packed down, ie compressed, with a press at a pressure between about 2 and about 12 bars, for a time ranging from 1 s to 20 s (Mäder Pressen) and/or with an isostatic press (ACB, France), at a 50 pressure between about 1,000 and about 1,500 bars, from about 1 minute to about 5 minutes, so as to reduce its thickness from about 1.5 to about 5 times, so as to get a final three dimensional collagen density ranging from about 50 mg/cm<sup>3</sup> to about 200 mg/cm<sup>3</sup>, either in a dry or wetted state.

Ten grams of the porous layer is further neutralized in one liter of isopropanol/ethanol (95/5, v/v) containing from about 20  $\mu$ mol to about 70  $\mu$ mol of sodium hydroxide. The alcohol mixture is further removed by several washings with acetone. The porous layer is then treated with a solution of HMDI at a 60 concentration of collagen of 50 grams per liter and at a concentration of HMDI ranging from about 0.1% to about 3% w/v. The mixture is let, under agitation, from 6 to 20 hrs, at ambient temperature, under mild agitation. The porous layer is then recovered by filtration removing acetone and is thoroughly washed with dry acetone to remove HMDI and acetone soluble HMDI byproducts. This HMDI cross-linking

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step can be repeated once or twice. At the end, the porous layer thus obtained is dried by removing the acetone residues.

A test has shown that the thus obtained porous layer is not substantially degraded by collagenase (Sigma), after 1 day in contact with collagenase, at +37° C. Generally, the degradation of the obtained HMDI cross-linked collagen by collagenase is less than 5% of its original amount whereas 65% or more of non cross-linked collagen is degraded in the very same conditions of the collagenase test.

### 3°) Coating of the Reinforcement Member

The coating reinforcement member is prepared as described in the Example 1, part 4°).

# 4°) Preparation of a Biosynthetic Mesh for Soft Tissue Repair

The biosynthetic mesh for soft tissue repair is prepared as described in Example 1, part 5°).

#### Example 7

# 1°) Preparation of a Biosynthetic Implant

A porous later has been prepared as described in the Example 6. The collagenic suspension is prepared by mixing the glutaraldehyde-cross-linked collagen and the oxidized collagen as obtained above in Example 6, at the following concentration:

- 1.125% m/v of glutaraldehyde-cross-linked collagen,
- 0.375% m/v of oxidized collagen.

Two hundreds and forty grams of this collagen suspension were poured within a 12 cm by 17 cm plastic box and freezedried for 48 hrs.

After the freeze-drying, the obtained porous layer was packed down, i.e., compressed, with the mechanical press (Mäger pressen) at 4 bars during two seconds and further packed down with the isostatic press (ACB, France) at 1,000 bars during 1 minute and then at 1,300 bars during 2 minutes.

The porous layer (50 grams) has been further treated with 1 liter of solution of HMDI at the concentration of 3% w/v in acetone, during 1 day. Then, the porous layer was extensively washed with acetone for removing HMDI and HMDI byproducts. The HMDI cross-linking step was repeated twice. At the end, the acetone was eliminated by evaporation.

The final thickness of the porous layer was about 1.5 mm with a corresponding three dimensional density of collagen at about 110 mg/cm<sup>3</sup>, either in a dry or wetted state.

The obtained porous layer was then further assembled with a coated reinforcement member, the reinforcement member being an isoelastic polyester mesh available under the tradename Parietex® C, from the company Sofradim, and showing a weight density of about 110 g/m². The Parietex® C mesh was coated as described in Example 1, part 4°).

#### 2°) Full Thickness Abdominal Wall Defect

A full thickness abdominal wall defect model has been designed in rat for the evaluation of the biosynthetic implants of the present Example. The surgery consisted in creating a 1.5×3.0 cm full defect of the lateral anterior parietal wall (fascia, muscle and peritoneum). The defect was covered with a 2.0×3.5 cm implant. On FIG. 10 are shown schematically the llinea alba 100, the skin 101 and the abdominal muscles 102 of a rat.

As a prophylactic measure the rats received a pre-operative subcutaneous injections of carprofene (Pfizer) and enrofloxacine (Bayer). The abdomen was clipped free of fur. The skin was scrubbed and painted with povidone iodine (Vetoquinol). With reference to FIG. 10, a medial incision of the abdominal skin 101 was performed. The left external surface of the abdominal wall was exposed. A full thickness 1.5×3 cm square defect 103 was created within the abdominal musculature 102. The peritoneum was removed together with the abdominal muscular layer. One defect 103 was created per animal. Each defect 103 was covered with an approximately 2×3.5 cm square implant 104 obtained in 1°) above, as depicted on FIG. 10. One product/implant 104 was implanted per animal, with smooth side on the flank of the abdominal cavity. The product was sutured using absorbable sutures 105 (such as Monocryl 4-0, Ethicon, France). The skin 101 was closed by continuous intracutaneous absorbable sutures (Mersilk 1 4Ph., Ethicon, France). A dressing was applied on the abdomen.

Macroscopic and microscopic observations of the implants were made at 2 weeks and 4 weeks. They showed the nice integration of the implant, fully covering the defect, starting to be cell colonized in its full thickness. No obvious signs of herniation, in particular at the edges of the implant in the abdominal wall were noticed. No significant degradation of the implant, in particular the mesh itself and the porous layer, was as well observed, even after 4 weeks,

It was noticed, even after 4 weeks of implantation, that the implant was not delaminated, for example having the mesh fully detached from the porous layer.

#### Example 8

The implant of Example 7 was repeated, wherein the polyester Parietex® C mesh was replaced by an isoelastic polypropylene mesh available under the tradename Parietene® PPL from the company Sofradim and having a weight density of about  $40~{\rm g/m^2}$ .

# Example 9

The implant of Example 7 was repeated, wherein the polyester Parietex® C mesh was replaced by an isoelastic polyester mesh available under the tradename Parietex® CL from the company Sofradim and having a weight density of about  $^{45}$  60 g/m².

#### Example 10

# 1°) Preparation of Porous Layers

The porous layers have been prepared as described in the Example 6. The collagenic suspension is prepared by mixing the glutaraldehyde-cross-linked collagen and the oxidized collagen obtained in Example 6 above, at the following concentrations:

Products A & B

1.125% m/v of glutaraldehyde-cross-linked collagen,

0.375% m/v of oxidized collagen.

A first product, the product A, was prepared by pouring one 60 hundred and eighty grams of this collagen suspension within a 12 cm by 17 cm plastic box and freeze-dried for 48 hrs. After the freeze-drying, the obtained porous layer was packed down with the isostatic press (ACB, France) at 1,000 bars during 1 minute and then at 1,300 bars during 2 minutes. It has 65 been packed in sealed aluminium/PET pouches, and further stored in an oven, at +103° C., during 1 day. It was then

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gamma-irradiated at a dose between 25 and 45 kGy and further heated at +50° C., during 2 days.

The final thickness of the porous layer was about 1.1 mm with a corresponding three dimensional density of collagen at about 135 mg/cm<sup>3</sup>, after its soaking in water during 24 hrs, at ambient temperature, i.e., in the wetted state of the porous layer.

A second product, the product B, was prepared by pouring two hundreds and forty grams of this collagen suspension within a 12 cm by 17 cm plastic box and freeze-dried for 48 hrs. After the freeze-drying, the obtained porous layer was packed down with the isostatic press (ACB, France) at 1,000 bars during 1 minute and then at 1,300 bars during 2 minutes. About 50 grams of the thus obtained porous layer has been further treated with 1 liter of solution of HMDI at the concentration of 3% w/v in acetone, during 1 day. Then, the porous layer was extensively washed with acetone for removing HMDI and HMDI byproducts. At the end, the acetone was eliminated by evaporation. It has been packed in sealed aluminium/PET pouches, and further stored in an oven, at +103° C., during 1 day. It was then gamma-irradiated at a dose between 25 and 45 kGy and further heated at +50° C., during 2 days.

The final thickness of the porous layer was about 0.9 mm with a corresponding three dimensional density of collagen at about 125 mg/cm<sup>3</sup>, after its soaking in water during 24 hrs, at ambient temperature, ie in a wetted state of the porous layer.

### 2°) Sub-Cutaneous Implantation of the Porous Layers A & B, in Rats

The sterile porous layers A & B, obtained as described above, were available as 10×20 mm pieces which were subsutaneously implanted in rats. The rats were sacrificed at 2, 4 and 12 weeks, after the implantation of the porous layers (three animals per time period with three sites per article and per time period). Macroscopic and histological observations have been performed for the evaluation of the implanted porous layers, in particular, for the evaluation of the local tolerance, the cellular or tissular colonization and implant degradation.

The study was based on the ISO10993 standard: Biological evaluation of medical devices, Part 6 (2007): Tests for local effects after implantation.

There was no evidence of morbidity or mortality during the study. No clinical abnormalities were detected during the course of the study. The porous layers A & B behaved in a similar way. At 2 weeks, the products were moderately integrated into the surrounding tissues with a moderate grade of cells and tissue colonization. A moderate grade of inflammation reaction was observed and products were slightly degraded by phagocytic cells. Very similar findings were observed at 4 weeks as compared to the 2 week time period with no fibrin and collagen deposit within the products, even if the product A showed better signs of colonization and integration. No significant increase of biodegradation was observed. At 12 weeks, very similar findings were also observed as compared to the previous time points (2 & 4 weeks), but with better tissue integration and cell colonization within the full thickness of the porous layers. The porous layers were slightly degraded with better resistance to degradation for the porous layer A.

At 2, 4 & 12 weeks, the initial thickness of the porous layers—as determined in a wet state, after the soaking of porous layers in water for 24 hours, at ambient temperature (18-25° C.), during 1 day—was not substantially reduced.

In conclusion, the local tolerance was good for the porous layers A& B with extended degradation times and with satisfying signs of cell colonization and integration of the porous layers.

It will be understood that various modifications may be 5 made to the embodiments disclosed herein. Therefore, the above description should not be construed as limiting, but merely as an exemplification of illustrative embodiments. Those skilled in the art will envision other modifications within the scope and spirit of the present disclosure. Such 10 modifications and variations are intended to come within the scope of the following claims.

What is claimed is:

- 1. An implant comprising:
- a porous layer comprising collagen;
- a non-porous layer comprising a collagenic constituent, the non-porous layer being joined to the porous layer; and
- a coated reinforcement member embedded into the nonporous layer,
- wherein the porous layer is compressed and has a three 20 dimensional density ranging from about 20 mg collagen/cm³ to about 200 mg collagen/cm³.
- 2. The implant of claim 1, wherein the porous layer has a three dimensional density ranging from about 50 mg collagen/cm<sup>3</sup> to about 150 mg collagen/cm<sup>3</sup>.
- 3. The implant of claim 1, wherein the collagen of the porous layer comprises oxidized collagen.
- **4**. The implant of claim **3**, wherein the collagen of the porous layer further comprises a collagen crosslinked with a compound selected from glutaraldehyde, hexamethylene 30 diisocyanate (HMDI) and mixtures thereof.

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- 5. The implant of claim 4, wherein the compound is glut-araldehyde.
- **6**. The implant of claim **4**, wherein the compound is hexamethylene diisocyanate.
- 7. The implant of claim 3, wherein the porous layer further comprises a chitosan.
- 8. The implant of claim 1, wherein the collagenic constituent of the non-porous layer is oxidized collagen.
- 9. The implant of claim 1, wherein the reinforcement member is a mesh having a weight density less than 100 g/m<sup>2</sup>.
  - 10. The implant of claim 9, wherein the mesh is isoelastic.
- 11. The implant of claim 1, wherein the reinforcement member is a mesh having a weight density greater than 50  $g/m^2$ .
- 12. The implant of claim 11, wherein the mesh is asymmetric.
- 13. The implant of claim 11, wherein the mesh is coated with a collagen crosslinked with a compound selected from glutaraldehyde, hexamethylene diisocyanate (HMDI) and mixtures thereof.
- **14**. The implant of claim **11**, wherein the mesh is coated with a mixture of oxidized collagen and chitosan.
- 15. The implant of claim 1, further comprising an additional non-porous layer.
- 16. The implant of claim 15, wherein the additional nonporous layer comprises an additional reinforcement member embedded therein.
- 17. The implant of claim 1, wherein the porous layer is biodegradable.

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